PCT

WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification ⁶:
 C12N 15/53, 9/02, 15/80, D21C 5/00

C12N 15/53, 9/02, 15/80, D21C 5/00, A61K 7/06, C12P 7/22, C12N 1/19, C09B 69/10 // (C12N 1/19, C12R 1:66) (11) International Publication Number:

WO 95/07988

(43) International Publication Date:

23 March 1995 (23.03.95)

(21) International Application Number:

PCT/US94/10264

A1

(22) International Filing Date:

13 September 1994 (13.09.94)

(30) Priority Data:

 08/122,230
 17 September 1993 (17.09.93)
 US

 08/122,827
 17 September 1993 (17.09.93)
 US

 08/162,827
 3 December 1993 (03.12.93)
 US

 08/172,331
 22 December 1993 (22.12.93)
 US

(71) Applicants: NOVO NORDISK A/S [DK/DK]; Novo Allé, DK-2880 Bagsværd (DK). NOVO NORDISK BIOTECH, INC. [US/US]; 1445 Drew Avenue, Davis, CA 95616-4880 (US).

(72) Inventors: WAHLEITHNER, Jill, Angela; 1718 Tea Place, Davis, CA 95616 (US). CHRISTENSEN, Bjærn, Eggert; Dronninggaards A11 32, DK-2840 Holte (DK). SCHNEIDER, Palle; Rydtoften 43, DK-2750 Ballerup (DK).

(74) Agents: ZELSON, Steve, T. et al.; Novo Nordisk of North America, Inc., Suite 6400, 405 Lexington Avenue, New York, NY 10174 (US). (81) Designated States: AM, AU, BB, BG, BR, BY, CA, CN, CZ, EE, FI, GE, HU, JP, KE, KG, KP, KR, KZ, LK, LR, LT, LV, MD, MG, MN, NO, NZ, PL, RO, RU, SD, SI, SK, TJ, TT, UA, UZ, VN, European patent (AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG), ARIPO patent (KE, MW, SD).

Published

With international search report.

Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments.

(54) Title: PURIFIED PH NEUTRAL RHIZOCTONIA LACCASES AND NUCLEIC ACIDS ENCODING SAME

(57) Abstract

The present invention relates to isolated nucleic acid fragments containing a sequence encoding a Rhizoctonia solani laccase having optimum activity at a neutral or basic pH, and the laccase proteins encoded thereby.

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AT	Austria	GB	United Kingdom	MR	Mauritania
ΑŪ	Australia	GE	Georgia	MW	Malawi
BB	Barbados	GN	Guinea	NE	Niger
BE	Belghun	GR	Greece	NL	Netherlands
BF	Burkina Paso	HU	Hungary	NO	Norway
BG	Bulgaria	DE.	Ireland	NZ	New Zealand
_	Benin	<u> </u>	<u>Liely</u>	PL.	Poland
BJ	- · 	JP	Japan	PT	Portugal
BR	Brazil	KE	Kenya	RO	Romania
BY	Belarus	KG	Kyrgystan	RU	Russian Pederation
CA	Canada		Democratic People's Republic	SD	Sudan
CF	Central African Republic	KP	• •	SE	Sweden
CG	Congo		of Korea	-	
CH	Switzerland	KR	Republic of Korea	SI	Slovenia
CI	Côte d'Ivoire	KZ	Kazakhstan	SK	Slovakia
CM	Cameroos	LI	Liechtenstein	SN	Senegal
CN	China	LK	Sri Lanka	TD	Chad
CS	Czechoslovakia	LU	Luxenbourg	TG	Togo
cz	Czech Republic	LV	Latvia	TJ	Tajikistan
DE	Germany	MC	Monaco	77	Trinidad and Tobago
DK	Denmark	MD	Republic of Moldova	UA	Ukraine
		MG	Madagascar	US	United States of America
ES	Spain		•	UZ	Uzbekistan
FI	Finland	ML	Mali	VN	Vict Nam
FR	Prance	MN	Mongolia	VN.	AND LIGHT
GA	Gabon				

PURIFIED PH NEUTRAL REIZOCTONIA LACCASES AND NUCLEIC ACIDS ENCODING SAME

5

10

15

Related Applications

This application is a continuation-in-part of copending U.S. Serial Nos. 08/122,230, 08/122,827, and 08/162,827, the contents of which are incorporated by reference in their entirety.

Field of the Invention

The present invention relates to isolated nucleic acid fragments encoding a fungal oxidoreductase enzyme and the purified enzymes produced thereby. More particularly, the invention relates to nucleic acid fragments encoding a phenol oxidase, specifically a laccase, which functions at a neutral pH.

20 Background of the Invention

Laccases (benzenediol:oxygen oxidoreductases) are multi-copper containing enzymes that catalyze the oxidation of phenolics. Laccase-mediated oxidations result in the production of aryloxy-radical intermediates from suitable phenolic substrate; the ultimate coupling of the intermediates so produced provides a combination of dimeric, oligomeric, and polymeric reaction products. Such reactions are important in nature in biosynthetic pathways which lead to the formation of melanin, alkaloids, toxins, lignins, and humic acids. Laccases are produced by a wide variety of fungi, including ascomycetes such as Aspergillus, Neurospora, and Podospora, the deuteromycete Botrytis, and

basidiomycetes such as Collybia, Fomes, Lentinus, Pleurotus, Trametes, and perfect forms of Rhizoctonia. Laccase exhibits a wide range of substrate specificity, and each different fungal laccase usually differs only quantitatively from others in its ability to oxidize phenolic substrates. Because of the substrate diversity, laccases generally have found many potential industrial applications. Among these are lignin modification, paper strengthening, dye transfer inhibition in detergents, phenol polymerization, juice manufacture, phenol resin production, and waste water treatment.

Although the catalytic capabilities are similar, laccases made by different fungal species do have different temperature and pH optima, and these may also differ 15 depending on the specific substrate. A number of these fungal laccases have been isolated, and the genes for several of these have been cloned. For example, Choi et al. (Mol. Plant-Microbe Interactions 5: 119-128, 1992) describe the molecular characterization and cloning of the gene encoding the laccase of the chestnut blight fungus, Cryphonectria parasitica. Kojima et al. (J. Biol. Chem. 15224-15230, 1990; JP 2-238885) provide a description 265: of two allelic forms of the laccase of the white-rot basidiomycete Coriolus hirsutus. Germann and Lerch 25 (Experientia <u>41</u>: 801,1985; PNAS USA <u>83</u>: 8854-8858, 1986) have reported the cloning and partial sequencing of the Neurospora crassa laccase gene. Saloheimo et al. (J. Gen. Microbiol. 137: 1537-1544, 1985; WO 92/01046) have disclosed a structural analysis of the laccase gene from the However, virtually all of the 30 fungus Phlebia radiata. known fungal laccases function best at acidic pHs (e.g., between pH 3.0 and 6.0), and are typically inactive at

neutral or basic pHs. Since a number of the aforestated potential industrial methods are preferentially conducted at neutral or basic pH, most fungal laccases perform poorly in such methods. Thus, the available <u>fungal laccases</u> are inadequate for application in a number of important commercial methods.

An exception to this rule is the extracellular laccase produced by certain species of Rhizoctonia. Bollag et al. have reported a laccase with a pH optimum of about 7.0 10 produced by Rhizoctonia praticola. A laccase of this type would be far more useful in industrial methods requiring neutral pH than previously known laccases. However, the R. praticola enzyme was neither purified nor further characterized, nor, to date, has any other laccase having 15 this trait been purified or characterized. Moreover, although other laccase genes have been isolated, as described above, these have been genes encoding enzymes which function best at acidic pH. Recombinant production and commercially adequate yields of a pH neutral or basic 20 laccase have thus been unattainable due to the fact that neither the enzyme per se nor the laccase gene encoding such an enzyme has previously been isolated and/or purified and sequenced. The present invention now provides a solution to each of these problems.

25

Summary of the Invention

The present invention relates to an isolated nucleic acid fragment comprising a nucleic acid sequence encoding a Rhizoctonia laccase which functions optimally at a pH between 6.0 to 8.5. By "functioning optimally" is meant that the enzyme exhibits significant(i.e., at least about 30% of maximum, preferably at least about 50%, and most

preferably from 50% to maximum) activity within the pH range of between about 6.0-8.5, as determined by activity in one or more standard laccase assays for substrates such as the syringaldazine, ABTS, 2.6-dimethoxyphenol, or 4

5 antiaminopyrine + N-ethyl-N-sulfobutyl-m-toluidine. A preferred substrate for the laccases of the present invention is syringaldazine. In a preferred embodiment, the laccase is a Rhizoctonia solani laccase. The invention also relates to a substantially pure laccase encoded by the novel nucleic acid sequence. By "substantially pure" is meant a laccase which is essentially (i.e., ≥90%) free of other non-laccase proteins.

In order to facilitate production of the novel laccase, the invention also provides vectors and host cells

comprising the claimed nucleic acid fragment, which vectors and host cells are useful in recombinant production of the laccase. The nucleic acid fragment is operably linked to transcription and translation signals capable of directing expression of the laccase protein in the host cell of

choice. A preferred host cell is a fungal cell, most preferably of the genus Aspergillus. Recombinant production of the laccase of the invention is achieved by culturing a host cell transformed or transfected with the nucleic acid fragment of the invention, or progeny thereof, under

conditions suitable for expression of the laccase protein, and recovering the laccase protein from the culture.

The laccases of the present invention are useful in a number of industrial processes in which oxidation of phenolics is required. These processes include lignin manipulation, juice manufacture, phenol polymerization and phenol resin production. In a preferred embodiment, the

enzyme of the invention is used in a process requiring a neutral or somewhat basic pH for greatest efficiency.

Brief Description of the Figures

Figure 1 illustrates the nucleotide and amino acid sequence of RSlac1. Lower case letters in the nucleotide sequence indicate the position of introns.

Figure 2 illustrates the nucleotide and amino acid sequence of RS1ac2. Lower case letters in the nucleotide sequence indicate the position of introns.

Figure 3 illustrates a restriction map of the plasmid pMWR-1.

Figure 4 illustrates the nucleotide and amino acid sequence of the translated region of RSlac3.

Figure 5 illustrates the syringaldazine oxidase activity of RSlac1 (90mM buffer, 20 μM syringaldazine, 20°C).

Figure 6 illustrates the syringaldazine oxidase activity of RSlac2 (93mM buffer, 20 µM syringaldazine, 20 20°C).

Detailed Description of the Invention

Certain species of the genus *Rhizoctonia* have been reported as producing laccase; therefore, an initial search focused on identifying the presence of these enzymes in various *Rhizoctonia solani* isolates. Samples are cultured and the supernatants periodically analyzed for the presence of laccase by the ABTS method, described below. Laccase is observed in all the *Rhizoctonia* cultures. Harvested laccases are electrophoretically separated and stained with ABTS. One isolate, RS22, produces a laccase with a basic pI, and is selected for further study.

The remaining studies focus on purification and characterization of the enzyme from RS22. Briefly, the fermentation broth is filtered and concentrated by UF with a membrane cut off of about 10,000. A first ion exchange chromatography step is conducted at pH 4.5 in acetate buffer, with step elution using NaCl. The eluate is then ultrafiltered and rechromatographed, and eluted with a NaCl gradient. Active fractions are pooled for further study.

The intact protein thus isolated and purified

(hereinafter referred to as RSlac3) is first subjected to

partial sequencing, and the N-terminal sequence obtained is

as follows:

AVRNYKFDIKNVNVAPDGFQRPIVSV (SEQ. ID. NO.: 5)

The protein is further subjected to digestion with a lysine- or glutamic-acid specific protease, and additional peptides obtained from the protein have the following sequences, which can be aligned with sequences in *Coriolus hirsutus*:

Peptide 1:

20 SQYVDGLRGPLVIYDPDDDH (SEQ. ID. NO: 6)

Peptide 2:

GLALVFAEAPSQIRQGVQSVQPDDA (SEQ. ID. NO.: 7)

Peptide 3:

SRYBVBBASTVVMLEBWYHTPAXVLE (SEQ. ID. NO. 8)

25 Peptide 4:

30

SLGPTPNYVNPXIRDVVRVGGTTVV (SEQ. ID. NO. 9)
The following peptides are also found, but do not correspond to *Coriolus* sequences

Peptide 5:

IRYVGGPAVX(N?)RSVI (SEQ. ID. NO.: 10)

Peptide 6:

ILANPA (SEQ. ID. NO.: 11)

PCT/US94/10264 **WO 95/07988**

Peptide 7:

15

25

YEAPSLPT (SEQ. ID. NO.: 12)

In the above sequences, B designates a residue which is either aspartic acid or asparagine, and X designates 5 unidentified residues.

In order to initiate screening for a Rhizoctonia laccase gene, an R. solani genomic library is prepared. Total DNA is partially digested with restriction enzyme Sau3A, and electrophoresed in an agarose gel to isolate DNA 10 fragments between 8 and 21 kb in size. The fractionated fragments are ligated to λ phage EMBL3 arms with BamHI ends, and the resulting phage packaged in vitro. These phage are used as a library to create a library of 170,000 plagues in E. coli and amplified 100-fold for future use.

In order to develop probes for isolation of the R. solani laccase gene, the protein sequences of five known laccases are analyzed to determine consensus sequences, and two degenerate oligonucleotides constructed based on observed consensus sequences (Choi et al. supra; Germann and 20 Lerch, supra; Saloheimo et al, supra, Kojima et al, supra). These oligos are mixed with R. solani genomic DNA and a DNA fragment of 220 nucleotide fragment is amplified using a tag polymerase chain reaction (PCR). The 220-nucleotide fragment is then cloned into plasmid vector.

The PCR fragment is used as a probe to screen 25,000 plaques from the amplified genomic library. Positive clones from this screen fall into two classes that are subsequently shown, by DNA sequence analysis, to code for two different laccase genes, RSlac1 and RSlac2. The nucleotide sequence 30 for each of these genes (SEQ ID. NOS.: 1 and 3), and the predicted amino acid sequence for each protein (SEQ. ID. NOS.: 2 and 4), are presented in, respectively, Figures 1

and 2. The homology between the two sequences is approximately 63%. Compared to known laccase sequences from Coriolus hirsutus, Phlebia radiata, Aspergillus nidulans, Cryphonectria parasitica, and Neurospora crassa, the RS laccases show between about 30-40% homology. Each of the two coding sequences is cloned into an expression vector operably linked to Aspergillus oryzae taka-amylase transcription and translation signals (See Figure 3). Each of the two laccase expression vectors is transformed into an Aspergillus oryzae and Aspergillus niger host cell, and the host cells screened for the presence of laccase.

For isolation of the RSlac3 gene, polyA RNA is purified from R. solani mycelia grown in the presence of anisidine. The RNA is used as a template for cDNA synthesis. The cDNA 15 is fractionated and fragments between 1.7-3.5 kb collected, and a cDNA library created by cloning the fractionated DNA into a yeast vector. 3000 transformants from this library are screened on ABTS. After 24 hours, a single colony appears positive. The plasmid from the colony is isolated 20 and the insert sequenced. Portions of the predicted amino acid sequence correspond with the sequences of the fragments obtained from RS 22, described supra. The complete nucleotide and amino acid sequences are depicted in Figure 4, and in SEQ. ID. NOS.: 13 and 14, respectively. RSlac3 25 shows 48% homology with RSlac1 and 50% homology with RSlac2. RSlac3 also shows 48% homology with the Coriolus hirsutus laccase gene.

According to the invention, a *Rhizoctonia* gene encoding a pH neutral or basic laccase can be obtained by 30 methods described above, or any alternative methods known in the art, using the information provided herein. The gene can be expressed, in active form, using an expression

vector. A useful expression vector contains an element that permits stable integration of the vector into the host cell genome or autonomous replication of the vector in a host cell independent of the genome of the host cell, and 5 preferably one or more phenotypic markers which permit easy selection of transformed host cells. The expression vector may also include control sequences encoding a promoter, ribosome binding site, translation initiation signal, and, optionally, a repressor gene or various activator genes. To 10 permit the secretion of the expressed protein, nucleotides encoding a signal sequence may be inserted prior to the coding sequence of the gene. For expression under the direction of control sequences, a laccase gene to be treated according to the invention is operably linked to the 15 control sequences in the proper reading frame. Promoter sequences that can be incorporated into plasmid vectors, and which can direct the transcription of the laccase gene, include but are not limited to the prokaryotic ß-lactamase promoter (Villa-Kamaroff, et al., 1978, Proc. Natl. Acad. 20 Sci. U.S.A. 75:3727-3731) and the tac promoter (DeBoer, et al., 1983, Proc. Natl. Acad. Sci. U.S.A. 80:21-25). Further references can also be found in "Useful proteins from recombinant bacteria* in Scientific American, 1980, 242:74-94; and in Sambrook et al., Molecular Cloning, 1989.

25

The expression vector carrying the DNA construct of the invention may be any vector which may conveniently be subjected to recombinant DNA procedures, and the choice of vector will typically depend on the host cell into which it is to be introduced. Thus, the vector may be an autonomously replicating vector, i.e. a vector which exists as an extrachromosomal entity, the replication of which is

independent of chromosomal replication, e.g. a plasmid, or an extrachromosomal element, minichromosome or an artificial chromosome. Alternatively, the vector may be one which, when introduced into a host cell, is integrated into the host cell genome and replicated together with the chromosome(s) into which it has been integrated.

In the vector, the DNA sequence should be operably connected to a suitable promoter sequence. The promoter may 10 be any DNA sequence which shows transcriptional activity in the host cell of choice and may be derived from genes encoding proteins either homologous or heterologous to the host cell. Examples of suitable promoters for directing the transcription of the DNA construct of the invention, 15 especially in a bacterial host, are the promoter of the lac operon of E.coli, the Streptomyces coelicolor agarase gene dagA promoters, the promoters of the Bacillus licheniformis α-amylase gene (amyL), the promoters of the Bacillus stearothermophilus maltogenic amylase gene (amyM), the 20 promoters of the Bacillus amyloliquefaciens α-amylase (amyQ), or the promoters of the Bacillus subtilis xylA and xylB genes. In a yeast host, a useful promoter is the eno-1 promoter. For transcription in a fungal host, examples of useful promoters are those derived from the gene encoding A. 25 oryzae TAKA amylase, Rhizomucor miehei aspartic proteinase, A. niger neutral α -amylase, A. niger acid stable α -amylase, A. niger or A. awamsii glucoamylase (gluA), Rhizomucor miehei lipase, A. oryzae alkaline protease, A. oryzae triose phosphate isomerase or A. nidulans acetamidase. Preferred 30 are the TAKA-amylase and gluA promoters.

The expression vector of the invention may also comprise a suitable transcription terminator and, in eukaryotes, polyadenylation sequences operably connected to the DNA sequence encoding the laccase of the invention.

5 Termination and polyadenylation sequences may suitably be derived from the same sources as the promoter. The vector may further comprise a DNA sequence enabling the vector to replicate in the host cell in question. Examples of such sequences are the origins of replication of plasmids pUC19, pACYC177, pUB110, pE194, pAMB1 and pIJ702.

The vector may also comprise a selectable marker, e.g. a gene the product of which complements a defect in the host cell, such as the dal genes from B.subtilis or B.li
15 cheniformis, or one which confers antibiotic resistance such as ampicillin, kanamycin, chloramphenicol or tetracycline resistance. Examples of Aspergillus selection markers include amds, pyrg, argB, niaD and sC, a marker giving rise to hygromycin resistance. Preferred for use in an

20 Aspergillus host cell are the amds and pyrg markers of A. nidulans or A. oryzae. A frequently used mammalian marker is the dihydrofolate reductase (DHFR) gene. Furthermore, selection may be accomplished by co-transformation, e.g. as described in WO 91/17243.

25

It is generally preferred that the expression is extracellular. The laccases of the present invention may thus comprise a preregion permitting secretion of the expressed protein into the culture medium. If desirable, this preregion may be native to the laccase of the invention or substituted with a different preregion or signal sequence, conveniently accomplished by substitution of the

DNA sequences encoding the respective preregions. For example, the preregion may be derived from a glucoamylase or an amylase gene from an Aspergillus species, an amylase gene from a Bacillus species, a lipase or proteinase gene from Rhizomucor miehei, the gene for the \alpha-factor from Saccharomyces cerevisiae or the calf prochymosin gene. Particularly preferred, when the host is a fungal cell, is the preregion for A. oryzae TAKA amylase, A. niger neutral amylase, the maltogenic amylase form Bacillus NCIB 11837, B. stearothermophilus \alpha-amylase, or Bacillus licheniformis subtilisin. An effective signal sequence is the A. oryzae TAKA amylase signal, the Rhizomucor miehei aspartic proteinase signal and the Rhizomucor miehei lipase signal.

The procedures used to ligate the DNA construct of the invention, the promoter, terminator and other elements, respectively, and to insert them into suitable vectors containing the information necessary for replication, are well known to persons skilled in the art (cf., for instance, Sambrook et al. Molecular Cloning, 1989).

The cell of the invention either comprising a DNA construct or an expression vector of the invention as defined above is advantageously used as a host cell in the recombinant production of a enzyme of the invention. The cell may be transformed with the DNA construct of the invention, conveniently by integrating the DNA construct in the host chromosome. This integration is generally considered to be an advantage as the DNA sequence is more likely to be stably maintained in the cell. Integration of the DNA constructs into the host chromosome may be performed

according to conventional methods, e.g. by homologous or heterologous recombination. Alternatively, the cell may be transformed with an expression vector as described above in connection with the different types of host cells.

5

The host cell may be selected from prokaryotic cells, such as bacterial cells. Examples of suitable bacteria are gram positive bacteria such as Bacillus subtilis, Bacillus licheniformis, Bacillus lentus, Bacillus brevis, Bacillus 10 stearothermophilus, Bacillus alkalophilus, Bacillus amyloliquefaciens, Bacillus coagulans, Bacillus circulans, Bacillus lautus, Bacillus megaterium, Bacillus thuringiensis, or Streptomyces lividans or Streptomyces murinus, or gram negative bacteria such as E.coli. The 15 transformation of the bacteria may for instance be effected by protoplast transformation or by using competent cells in a manner known per se.

The host cell may also be a eukaryote, such as mammalian cells, insect cells, plant cells or preferably

20 fungal cells, including yeast and filamentous fungi. For example, useful mammalian cells include CHO or COS cells. A yeast host cell may be selected from a species of Saccharomyces or Schizosaccharomyces, e.g. Saccharomyces cerevisiae. Useful filamentous fungi may selected from a

25 species of Aspergillus, e.g. Aspergillus oryzae or Aspergillus niger. Alternatively, a strain of a Fusarium species, e.g. F. oxysporum, can be used as a host cell. Fungal cells may be transformed by a process involving protoplast formation and transformation of the protoplasts followed by regeneration of the cell wall in a manner known per se. A suitable procedure for transformation of Aspergillus host cells is described in EP 238 023. A suitable method of

transforming Fusarium species is described by Malardier et al., 1989.

The present invention thus provides a method of producing a recombinant laccase of the invention, which sethod comprises cultivating a host cell as described above under conditions conducive to the production of the enzyme and recovering the enzyme from the cells and/or culture medium. The medium used to cultivate the cells may be any conventional medium suitable for growing the host cell in question and obtaining expression of the laccase of the invention. Suitable media are available from commercial suppliers or may be prepared according to published formulae (e.g. in catalogues of the American Type Culture Collection).

The resulting enzyme may be recovered from the medium by conventional procedures including separating the cells from the medium by centrifugation or filtration, precipitating the proteinaceous components of the supernatant or filtrate by means of a salt, e.g. ammonium sulphate, followed by purification by a variety of chromatographic procedures, e.g. ion exchange chromatography, gel filtration chromatography, affinity chromatography, or the like. Preferably, the isolated protein is about 90% pure as determined by SDS-PAGE, purity being most important in food, juice or detergent applications.

In a particularly preferred embodiment, the expression of laccase is achieved in a fungal host cell, such as Aspergillus. As described in detail in the following examples, the laccase gene is ligated into a plasmid containing the Aspergillus oryzae TAKA α-amylase promoter, and the Aspergillus nidulans amdS selectable marker. Alternatively, the amdS may be on a separate plasmid and

used in co-transformation. The plasmid (or plasmids) is used to transform an Aspergillus species host cell, such as A. oryzae or A. niger in accordance with methods described in Yelton et al. (PNAS USA 81: 1470-1474,1984).

Those skilled in the art will recognize that the 5 invention is not limited to use of the nucleic acid fragments specifically disclosed herein, for example, in Figures 1 and 2. It will be apparent that the invention also encompasses those nucleotide sequences that encode the 10 same amino acid sequences as depicted in Figures 1, 2 and 3, but which differ from those specifically depicted nucleotide sequences by virtue of the degeneracy of the genetic code. In addition, the invention also encompasses other nucleotide fragments, and the proteins encoded thereby, which encode 15 laccase proteins having substantially the same pH optimum as those of Rhizoctonia solani, and which show a significant level of homology with the Rhizoctonia solani amino acid sequence. For example, the present data show that more than one species of Rhizoctonia produces a laccase with the 20 desired pH profile; it is therefore expected that other Rhizoctonia species also produce similar laccases and therefore, using the technology described herein, can be used as a source for genes within the scope of the claimed invention. As also shown in the present examples, not only 25 is there more than one nucleotide and amino acid sequence that encodes a laccase with the required characteristics, there is also considerable variation tolerated within the sequence while still producing a functional enzyme. Therefore, the invention also encompasses any variant 30 nucleotide sequence, and the protein encoded thereby, which protein retains at least about an 80% homology with one or the other of the amino acid sequences depicted in Figures 1,

2 and 3, and retains both the laccase and pH optimum activity of the sequences described herein. In particular, variants which retain a high level(i.e., ≥ 80%) of homology at highly conserved regions of the Rhizoctonia laccase are contemplated. Such regions are identified as residues 458-469 in RSLAC1, and 478-489 in RSLAC2; and residues 131-144 in RSLACI and 132-145 in RSLAC2.

Useful variants within the categories defined above include, for example, ones in which conservative amino acid 10 substitutions have been made, which substitutions do not significantly affect the activity of the protein. By conservative substitution is meant that amino acids of the same class may be substituted by any other of that class. For example, the nonpolar aliphatic residues Ala, Val, Leu, 15 and Ile may be interchanged, as may be the basic residues Lys and Arg, or the acidic residues Asp and Glu. Similarly, Ser and Thr are conservative substitutions for each other, as are Asn and Gln. It will be apparent to the skilled artisan that such substitutions can be made outside the 20 regions critical to the function of the molecule and still result in an active enzyme. Retention of the desired activity can readily be determined by conducting a standard ABTS oxidation method in 0.1M sodium phosphate at pH 7.0.

The protein can be used in number of different

industrial processes; although the enzyme is also functional
to some extent at lower pH, the R. solani laccase is most
beneficially used in processes that are usually conducted at
a neutral or alkaline pH, since other laccases are not
active in this pH range. These processes include

polymerization of lignin, both Kraft and lignosulfates, in
solution, in order to produce a lignin with a higher
molecular weight. A neutral/alkaline laccase is a

particular advantage in that Kraft lignin is more soluble at higher pHs. Such methods are described in, for example, Jin et al., Holzforschung 45(6): 467-468, 1991; US Patent No. 4.432.921; EP 0 275 544; PCT/DK93/00217, 1992.

5 The laccase of the present invention can also be used for in-situ depolymerization of lignin in Kraft pulp, thereby producing a pulp with lower lignin content. This use of laccase is an improvement over the current use of chlorine for depolymerization of lignin, which leads to the production of chlorinated aromatic compounds, which are an environmentally undesirable by-product of paper mills. Such uses are described in, for example, Current opinion in Biotechnology 3: 261-266, 1992; J. Biotechnol. 25: 333-339, 1992; Hiroi et al., Svensk papperstidning 5: 162-166, 1976.

15 Since the environment in a paper mill is typically alkaline, the present laccase is more useful for this purpose than other known laccases, which function best under acidic conditions.

Oxidation of dyes and other chromophoric compounds
leads to decolorization of the compounds. Laccase can be
used for this purpose, which can be particularly
advantageous in a situation in which a dye transfer between
fabrics is undesirable, e.g., in the textile industry and in
the detergent industry. Methods for dye transfer inhibition
and dye oxidation can be found in WO 92/01406, WO 92/18683,
EP 0495836 and Calvo, Mededelingen van de Faculteit
Landbouw-wetenschappen/Rijiksuniversitet Gent.56: 1565-1567,
1991.

The present laccase can also be used for the
polymerization of phenolic compounds present in liquids. An
example of such utility is the treatment of juices, such as
apple juice, so that the laccase will accelerate a

precipitation of the phenolic compounds present in the juice, thereby producing a more stable juice. Such applications have been described in Stutz, Fruit processing 7/03, 248-252, 1993; Maier et al., Dt. Lebensmittel
rindschau 86(5): 137-142, 1990; Dietrich et al., Fluss. Obst 57(2): 67-73, 1990. The invention is further illustrated by the following non-limiting examples.

EXAMPLES

1. Purification and characterization of R. solani laccase

Individual isolates of R. solani cultured on potato dextrose agar (Difco) are examined for laccase enzyme formation by transferring a small piece of agar containing vigorous growth to 100 ml CFM (24.0 g potato dextrose broth, 3.0 g yeast extract, 1.0 ml Microelement solution

[0.80 g KH₂PO₄, 0.64 g CuSO₄·5H₂O, 0.11 g FeSO₄·7H₂O, 0.80 g MnCl₂·4H₂O, 0.15 g ZnSO₄·7H₂O, distilled water to 1000 ml], distilled water to 1000 ml) in a 500 ml shake flask. Incubation is at room temperature, at 200 rpm on an orbital shaker.

Samples are harvested at 50, 74, 122 and 170 hours, centrifuged and the clear supernatant analyzed for laccase with its ABTS (ABTS= 2,2'-azinobis (3 ethylbenzothiazoline-6-sulfonic acid). The analysis is carried out by adding 200 µl of 2mM ABTS in 0.1 M phosphate buffer, pH 7, and observing the change in absorbance at 418 nm after 30 minutes incubation at room temperature (approximately 23-25°C). This method is modified from a peroxidase analysis method described by Pütter and Becker (Peroxidases, in: Bergmeyer, H.U.(ed.), Methods of Enzymatic Analysis, 3rd ed., Vol.III, pp.286-293, 1983)

Each of the laccases harvested at 172 hours is electrophoretically separated and stained with ABTS as

chromogen. Several distinct patterns emerge; the strain RS 22 is shown to produce a laccase having a basic pI, and is chosen for further characterization.

Laccase activity is also determinable by the 5 syringaldazine method. Laccase catalyzes the oxidation of syringaldazine to tetramethoxy azo bis-methylene quinone under aerobic conditions, with a change of color from yellow to violet. 3000 µl of 25 mM acetate buffer (containing 10mg/l cuprisulfate, 5 H_2O) at pH 5.5, 30°C, is mixed in a 1 10 cm cuvette with 225 μ l 0.28 mM syringaldazine (5mg solubilized in 25 ml ethanol and adjusted to 50 ml with demineralized water). The mixture is then mixed with 100 μ l of a laccase dilution (diluted in acetate buffer so that the increase in absorbance (Δ OD) is within the range of 0.1-0.6). 15 The reaction mixture is placed in a 30°C thermostated spectrophotometer and the reaction is followed at 530 nm for 10 to 70 seconds from the addition of laccase. The activity of the enzyme is calculated as $\Delta OD/minute \times 0.677 \times dilution$ factor, and is expressed as LACU.

For purification of the Rhizoctonia laccase, 2.1 liter of culture medium with a LACU activity of 0.19 LACU/ml is filtered through a 10 μm filter and concentrated to 230 ml by ultrafiltration using a Filtron Minisette OMEGA membrane with a cutoff value of 10 kDa. The pH of the sample is 5.3 and the activity of the concentrated sample is determined to be 3.34 LACU/ml.

After pH adjustment to 4.5 and filtration due to slight precipitation, the sample is applied to a 40 ml S Sepharose Fast Flow column equilibrated with 20mM acetate buffer at pH 30 4.5 (buffer A). The column is washed in buffer A and eluted with buffer A containing 1 M NaCl. Active fractions are collected and pooled. This active pool is concentrated and

buffer exchanged to buffer A using an Amicon ultrafiltration unit equipped with a Diaflo YM10 membrane. This sample is rechromatographed on a 5 ml S Sepharose High Performance column using the method described above except that clution is carried out with a linear gradient over 30 column volumes from buffer A to buffer A containing 0.5 M NaCl. The fractions from this purification exhibiting highest activity are pooled. Approximately 45 mg laccase are obtained, when protein concentration is estimated by one absorption unit at A280 nm equal to lmg/ml. The protein is >90% pure as judged by SDS-PAGE. The molecular weight estimated by SDS-PAGE is approximately 67 kDa. The specific activity of the purified protein is 1 LACU/mg. The pH profile of the purified protein, using syringaldazine as substrate is show in Table 1, below.

Table 1.

	Ha	5	6	7	8
20	% activity	0.5	31	100	59

For sequencing of the protein, peptides are generated using wither a lysine-specific protease from Achromobacter (Achromobacter protease I) or a glutamic acid specific protease from Bacillus licheniformes. The peptides are purified by reverse phase HPLC employing linear gradients of 80% 2-propanol containing 0.08% aqueous TFA (solvent B) in 0.1% aqueous TFA (solvent A).

N-terminal amino acid sequence analysis of the intact 30 protein and of purified peptides are carried out in an Applied Biosystems 473A protein sequencer according to the manufacturer's instructions. Initial partial sequencing of

the isolated protein yields the following N-terminal sequence:

AVRNYKFDIKNVNVAPDGFQRPIVSV (SEQ. ID. NO.: 5)

The protein is then digested with either a lysine- or glutamic-acid specific protease, and following additional peptides identified. Peptides 1-4 can be aligned with sequences in the laccase of *Coriolus hirsutus*:

Peptide 1:

SOYVDGLRGPLVIYDPDDDH (SEQ. ID. NO: 6)

10 Peptide 2:

GLALVFAEAPSQIRQGVQSVQPDDA (SEQ. ID. NO.: 7)

Peptide 3:

SRYBVBBASTVVMLEBWYHTPAXVLE (SEQ. ID. NO. 8)

Peptide 4:

15 SLGPTPNYVNPXIRDVVRVGGTTVV (SEQ. ID. NO. 9)

Peptide 5:

IRYVGGPAVX(N?)RSVI (SEQ. ID. NO.: 10)

Peptide 6:

ILANPA (SEQ. ID. NO.: 11)

20 Peptide 7:

YEAPSLPT (SEQ. ID. NO.: 12)

An X in the above sequences designates an unidentified residue, and B represents a residue which is either aspartic acid or asparagine.

25

2. Isolation of R. solani laccase gene

A study of the known amino acid sequences of fungal laccases obtained from non-Rhizoctonia species (Choi et al., supra; German et al., supra; Saloheimo et al. supra; and Kojima et al, supra) is conducted to determine the presence of consensus sequences among them. Two regions of high identity, IHWHGFFQ and TFWYHSH, are found near the amino

terminal third of the protein. Based on these consensus sequences and the corresponding DNA sequences, three degenerate oligonucleotides, O-lac2 [TGG/AAAGACCATA/GGTGTCG/AGTA/G], its complement O-lac2r, and

[TGG/AAAGACCATA/GGTGTCG/AGTA/G], its complement O-lac21, and o-lac3[ATCCAT/CTGGCAT/CGGG/CA/TTCTTCCAG/A], are synthesized using an Applied Biosystems 394 DNA/RNA synthesizer.

The synthesized oligos are used in a polymerase chain reaction (PCR) to screen Rhizoctonia solani genomic DNA for a laccase gene or fragment thereof. For amplifications of genomic DNA, 0.5 µg of genomic DNA is incubated with 1µM of each primer, 200µM of dNTPs, and 1 U taq polymerase (Boehringer Mannheim) in [10 mM Tris-Cl, 1.5 mM MgCl₂, 50 mM KCl, 1 mg/ml gelatine;pH 8.3]. The reactions are incubated for 1x5 minutes at 95°C, 30x[1 minute at 95°C, 1 minute at 50-60°C, 1 minute at 72°C], and 1x5 minutes at 72°C. The PCR reactions amplify a DNA fragment of 220 nucleotides. The PCR product is cloned, according to manufacturer's directions, into the TA cloning vector (InVitrogen Corp.). Characterization of the PCR product by DNA sequencing of individual clones distinguishes two separate laccase genes designated RSlacl and RSlac2.

To prepare a R. solani genomic library, R. solani DNA is partially digested with restriction enzyme Sau3A, and electrophoresed through a 0.8% Sea Plaque Agarose (FMC Bioproducts) in a Tris/Acetate/EDTA buffer to isolate those DNA fragments between 8.0 an 21 kb in size. The gel fractionated fragments are further purified with Beta-Agarase(New England Biolabs) according to manufacturer's instruction, and then ligated to lambda phage EMBL3 arms with BamHI ends. The resulting phages are packaged in vitro using Gigapack II packaging extract(Stratagene). 25 ml of TB media+0.2% maltose and 10 MgSO4 is inoculated into a 50 μl

aliquot of an overnight culture of E. coli K802 (supE, hsdR, gal, metB) and incubated at 37°C with shaking until the A600=0.5. 25 µl of a 1:10 and 1:50 dilution of the packaged phage are mixed with 250 µl of the K802 cells, and incubated for 20 minutes at 37°C. To each dilution, 5 µl of melted top agar at 48°C are added. The mix is then plated onto prewarmed LB plates and incubated at 37°C for at least 12 hours. From these phage, a library of 170,000 plaques in E.coli K802 is created and amplified 100-fold for future use.

To screen for the laccase gene, 25,000 plaques from the amplified genomic library are plated onto NZY/agarose plates for plaque lifts using conventional methods. Filters are screened using the 220 nucleotide PCR fragment randomly

15 labelled to 5x108 cpm/µg as a probe. Filters are hybridized in 50% formamide, 6xSSC for 16 hours at 42°C and washed with 0.5xSSC, 0.1% SDS at 65°C. Positive clones are picked and rescreened using conventional methods. The nine positive clones identified fell into two classes that by DNA sequence analysis are shown to code for two different laccase genes, RSlac1 and RSlac2. The complete nucleotide sequence of each of these genes is determined using fluorescent nucleotides and an Applied Biosystems automatic DNA sequencer (Model 363A, version 1.2.0). The nucleotide and predicted amino acid sequences are depicted in Figures 1 and 2.

For isolation of RSlac3, poly A RNA purified from R.

solani mycelia grown in the presence of 1 mM anisidine is
used as a template for cDNA synthesis using standard
protocols. The cDNA is fractionated by electrophoresis
through a 0.8% agarose gel and DNA fragments between 1.7 and
3.5 kb in size are collected. A library is then created by
cloning the size-fractionated cDNA into the yeast expression

vector pYES2. 3000 yeast transformants from this library are plated initially on YNB (1.7 g yeast nitrogen base without amino acids, 5 g (NH₄)₂SO₄ per liter) with 2% glucose. After 4 days growth at 30 C, the resulting colonies are replica plated to YNB with 0.1% glucose, 2% galactose and 2mM ABTS [2,2'-azino-bis(3-ethylbenzthiazoline-6-sulfonic acid; Sigma # A-1888). After 24 hours of growth at 30 C a single colony has a light green halo which gradually turns a dark purple. The plasmid from this colony is isolated and the insert sequenced. The sequence of the translated portion of the RSlac3 gene and protein is shown in SEQ.ID NOS. 13 and 14, and in Figure 4. 3. Expression of laccase gene

, An

The plasmid pMWR-1 is a pUC derived vector containing
the TAKA amylase transcription regulation signals and the
TAKA amylase signal sequence. This plasmid is engineered
with a unique SfiI site at the signal sequence cleavage
site, and a 3' adjacent NsiI site such that these two
restriction enzymes can be used to introduce, in frame, a
foreign protein. Using a PCR reaction (conducted as
described above, but with 100 ng of the appropriate
linearized plasmid DNA as a template) and mutagenized
primers, an SfiI site is introduced at amino acid 12 and
amino acid 14 of RSlac1 and RSlac2, respectively, such that
the protein coding sequences are in frame with the TAKA
signal sequence. In addition, a PCR amplification is also
used to introduce a PstI site (CTGCAG) at the 3' end of
RSlac1 and an NsiI site (ATGCAT) at the 3' end of RSlac2.

To prepare for transformation, cells of Aspergillus oryzae are cultivated in YPG (1g/l yeast extract, 0.25 g K₂PO₄. 0.125 g/MgSO₄, 3.75 g glucose) at 34°C with 100-120rpm

for 16-20 hours, then collected by filtration with miracloth. Cells are washed with Mg solution (0.6M MgSO₄·7H₂O), then 2-6 g of cells are taken up in 10 ml MgF(1.2M MgSO₄·7H₂O, 10mM NaH₂PO₄·2H₂O;pH 5.8).To this is added 1 ml of Novozyme® 234 (120 mg/ml MgP), and the sample kept on ice for 5 minutes. One ml of BSA (12 mg/ml) is added, and the sample shaken gently at 34-37°C. Protoplasts are collected by filtration through miracloth, and overlain with 5 ml of ST (0.6 M Sorbitol, 100mM Tris; pH 7). The sample is spun at 2500 rpm for 15 minutes, and a band of protoplasts collected. Two volumes of STC (1.2M Sorbitol, 10 mM tris, 10 mM CaCl₂·2H₂O;pH 7.5) are added and the sample is spun at 2500 rpm for 5 minutes. The precipitate is washed twice with 5 ml of STC, and the protoplasts suspended in 0.5-lml of STC.

For the transformation process, the protoplast concentration is adjusted to 1-5x10 $^7/ml$. To 100 μl of protoplast solution is added a maximum of 10 µl of DNA solution (5-10 μ g of supercoiled DNA) and 0.2 ml of PEG 20 (60% PEG4000, 10mM Tris, 10mM CaCl₂·H₂O; pH 7.5), and the combination is mixed well. The sample is kept at room temperature for 25 minutes; then to it is added first 0.2 ml PEG, with mixing, the 0.85 ml PEG with mixing. The mixture is kept at room temperature for 20 minutes, then spun at 25 4000 rpm for 15 minutes. The precipitate is washed with 2 ml of STC by spinning at 2500 rpm for 10 minutes. protoplasts are resuspended in 0.2-0.5 ml of STC, and then spread on COVE plates. COVE medium (pH 7) contains 342.3 g/l sucrose, 25 g/l agar and a salt solution comprising 26 g/l 30 KCl, 26 g/l MgSO₄· H_2 O, 76 g/l K H_2 PO₄, and 50 ml/l of trace metals; the trace metals are 40 mg/l $NaB_4O_7 \cdot 10H_2O$, 400 mg/l

PCT/US94/10264

CuSO₄·5H₂O, 1200mg/l FeSO₄·7H₂O, 700mg/l MnSO₄·H₂O, 800mg/l Na₂MoO₂·2H₂O, 10 g/l ZnSO₄·7H₂O). After autoclaving, 10 ml/l of 1M filtrated acetamide and 5-10 ml of 3M CsCl are added to the solution. Transformants are selected by growth cells on COVE medium which contains acetamide as the carbon source.

The confirmation of laccase production in the samples is determined by the ABTS oxidation method as described above on Cove medium with 2 mM ABTS, at pH 5 and 7.3. Both RSlac1 and RSlac2 express laccase activity at pH 5 and pH 7, in contrast with a control laccase which shows substantially no activity at pH 7.3.

The products of the expression of each of RSlacl and RSlac2 are tested for oxidase activity at various pHs using syringaldazine as the substrate. The assay is conducted substantially as described above for the assay of the native protein, over pH range of 4-9. As shown in Figures 5 and 6, both laccases are active at pHs over pH 5, and RSlacl has particularly good activity at pHs over 6. The pattern of activity is generally comparable to that observed for the RSlac3 laccase isolated from RS 22 (see Table 1 above), with RSlac1 exhibiting the broadest range of activity.

Deposit of Biological Materials

The following biological materials have been deposited under the terms of the Budapest Treaty in the International Mycological Institute, Genetic Resource Reference Collection, located at Bakeham Lane, Egham, Surrey TW20 9TY and given the following accession number.

30 <u>Deposit</u>

Rhizoctonia solani RS22

Accession Number
IMI CC 358730

PCT/US94/10264

The following biological materials have been deposited under the terms of the Budapest Treaty with the Agricultural Research Service Patent Culture Collection, Northern Regional Research Center, 1815 University Street, Peoria,

5 Illinois, 61604 and given the following accession numbers.

<u>Deposit</u>

Accession Number

E. coli containing RSlacl fused to

NRRL B-21141

an α -amylase signal sequence

(EMCC 00844)

10

WO 95/07988

E. coli containing RSlac2 with an SfiI site insert (EMCC 00845)

NRRL B-21142

15 E. coli containing RSlac3
(EMCC 0088)

NRRL B-21156

PCT/US94/10264 WO 95/07988

SEQUENCE LISTING

(1) GENERAL INFORMATION:

- (i) APPLICANT:
 - (A) NAME: Novo Nordisk A/S
 - (B) STREET: Novo Alle
 - (C) CITY: Bagsværd
 - (D) COUNTRY: Denmark
 - (E) POSTAL CODE (ZIP): DK-2880 (F) TELEPHONE: +45 4444 8888

 - (G) TELEFAX: +45 4449 3256
 - (F) TELEX: 37304

(i) APPLICANT:

- (A) NAME: Novo Nordisk Biotech, Inc.
- (B) STREET: 1445 Drew Avenue
- (C) CITY: Davis, California
- (D) COUNTRY: United States of America
- (E) POSTAL CODE (ZIP): 95616-4880
- (F) TELEPHONE: (916) 757-8100
- (G) TELEFAX: (916) 758-0317
- (ii) TITLE OF INVENTION: PURIFIED PH NEUTRAL LACCASES AND NUCLEIC ACIDS ENCODING SAME
- (iii) NUMBER OF SEQUENCES: 14
- (iv) CORRESPONDENCE ADDRESS:
 - (A) ADDRESSEE: Novo Nordisk of North America, Inc.
 - (B) STREET: 405 Lexington Avenue, Suite 6400
 - (C) CITY: New York
 - (D) STATE: New York
 - (E) COUNTRY: USA
 - (F) ZIP: 10174-6401
 - (v) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Floppy disk
 - (B) COMPUTER: IBM PC compatible
 - (C) OPERATING SYSTEM: PC-DOS/MS-DOS
 - (D) SOFTWARE: PatentIn Release #1.0, Version #1.25
- (vi) CURRENT APPLICATION DATA:
 - (A) APPLICATION NUMBER: to be assigned
 - (B) FILING DATE: 13-SEP-1994
- (vii) PRIOR APPLICATION DATA:
 - (A) APPLICATION NUMBER: US 08/172,331
 - (B) FILING DATE: 22-DEC-1993
- (vii) PRIOR APPLICATION DATA:
 - (A) APPLICATION NUMBER: US 08/122,230
 - (B) FILING DATE: 17-SEP-1993
- (vii) PRIOR APPLICATION DATA:
 - (A) APPLICATION NUMBER: US 08/122,827
 - (B) FILING DATE: 17-SEP-1993
- (vii) PRIOR APPLICATION DATA:
 - (A) APPLICATION NUMBER: US 08/162,827
 - (B) FILING DATE: 03-DEC-1993
- (viii) ATTORNEY/AGENT INFORMATION:

 - (A) NAME: Lowney Dr., Karen A.(B) REGISTRATION NUMBER: 31,274
 - (C) REFERENCE/DOCKET NUMBER: 4052.204-WO

- (ix) TELECOMMUNICATION INFORMATION:
 - (A) TELEPHONE: 212-867-0123
 - (B) TELEFAX: 212-878-9655

(2) INFORMATION FOR SEQ ID NO:1:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 2838 base pairs

 - (B) TYPE: nucleic acid (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Rhizoctonia laccase
- (ix) FEATURE:
 - (A) NAME/KEY: intron
 - (B) LOCATION: 302..351
- (ix) FEATURE:
 - (A) NAME/KEY: intron
 - (B) LOCATION: 463..512
- (ix) FEATURE:
 - (A) NAME/KEY: intron
 - (B) LOCATION: 576..633
- (ix) FEATURE:

 - (A) NAME/KEY: intron
 (B) LOCATION: 760..818
- (ix) FEATURE:
 - (A) NAME/KEY: intron
 - (B) LOCATION: 822..877
- (ix) FEATURE:
 - (A) NAME/KEY: intron
 - (B) LOCATION: 1001..1054
- (ix) FEATURE:
 - (A) NAME/KEY: intron
 - (B) LOCATION: 1316..1372
- (ix) FEATURE:
 - (A) NAME/KEY: intron
 - (B) LOCATION: 1697..1754
- (ix) FEATURE:
 - (A) NAME/KEY: intron
 - (B) LOCATION: 1827..1880
- (ix) FEATURE:
 - (A) NAME/KEY: intron
 - (B) LOCATION: 1992..2051
- (ix) FEATURE:
 - (A) NAME/KEY: intron
 - (B) LOCATION: 2157..2206
- (ix) FEATURE:
 - (A) NAME/KEY: intron
 - (B) LOCATION: 2348..2404
- (ix) FEATURE:

NAME/KEY: LOCATION:	intron 24382498

(ix) FEATURE:

(A) NAME/KEY: CDS
(B) LOCATION: join(170..301, 352..462, 513..575, 634..759, 819
..821, 878..1000, 1055..1315, 1373..1696, 1755
..1826, 1881..1991, 2052..2156, 2207..2347, 2405
..2437, 2499..2621)

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

(XI) SEQUENCE DESCRIPTION: SEQ ID NO.I.				
AGCGTCACAC CAGACATCGG ATGAAAACGG AAAGTGTATG CGCCATTTGA CGTCTGCGGC	60			
AACCACTGTT CATCTCGCGA GCTAACATGG GCGACGTATA AGAAGAACGC GAGAATGGGC	120			
AGATTTCGAT ATCCCCTCTC GTCTCGGTTT TGGTCTCGGC TTGCCTCTA ATG GCG Met Ala 1	175			
CGC ACC ACT TTC CTT GTC TCG GTT TCG CTC TTT GTT TCC GCT GTT CTT Arg Thr Thr Phe Leu Val Ser Val Ser Leu Phe Val Ser Ala Val Leu 5 10 15	223			
GCG CGC ACC GTC GAG TAC GGC TTG AAG ATT AGT GAT GGG GAG ATA GCT Ala Arg Thr Val Glu Tyr Gly Leu Lys Ile Ser Asp Gly Glu Ile Ala 20 25 30	271			
CCT GAC GGT GTT AAG CGT AAT GCG ACT TTG GTACGCACTC CTTGTAATCC Pro Asp Gly Val Lys Arg Asn Ala Thr Leu 35 40	321			
AACAATTCAA GGTTTCTGAT GCTTGGTCAG GTA AAT GGA GGG TAT CCC GGT CCA Val Asn Gly Gly Tyr Pro Gly Pro 45 50	375			
CTC ATT TTT GCC AAC AAG GGG GAT ACT CTC AAA GTC AAG GTC CAA AAC Leu Ile Phe Ala Asn Lys Gly Asp Thr Leu Lys Val Lys Val Gln Asn 55 60 65	423			
AAG CTC ACG AAT CCT GAG ATG TAT CGC ACC ACT TCC ATC GTATGTTCGT Lys Leu Thr Asn Pro Glu Met Tyr Arg Thr Thr Ser Ile 70 75 80	472			
TCGATATCTA CTAATACATC CGTCGCTAAA TATCTTGTAG CAT TGG CAC GGT CTC His Trp His Gly Leu 85	527			
TTA CAA CAT AGA AAC GCC GAC GAC GGC GGT CCT TCG TTC GTC ACT CAG Leu Gln His Arg Asn Ala Asp Asp Gly Pro Ser Phe Val Thr Gln 90 95 100	575			
GTAGGATTCT GGAAGGTTGG CCTGAACTCT CTGTTAACCG ACAACCCGAT GTCACCAG	633			
TGC CCG ATT GTT CCA CGC GAG TCG TAT ACT TAC ACC ATA CCT CTG GAC Cys Pro Ile Val Pro Arg Glu Ser Tyr Thr Tyr Thr Ile Pro Leu Asp 105 110 115	681			
GAT CAA ACC GGA ACC TAT TGG TAC CAT AGC CAC TTG AGT TCG CAA TAC Asp Gln Thr Gly Thr Tyr Trp Tyr His Ser His Leu Ser Ser Gln Tyr 120 130	729			
GTT GAT GGT CTT CGA GGC CCG CTG GTA ATC GTGAGTATCT TGACTTGTCT Val Asp Gly Leu Arg Gly Pro Leu Val Ile 135 140	779			

(
ACTGAAGGCA ACGAGACTAA AACAAGCGTC GATTCACAG TAT GTTCGTCTCC Tyr 145	831
CCTTTATTTA GCTCTGGATC TTCATTTCTC ACGTAATACA TGATAG GAT CCC AAG Asp Pro Lys	886
GAT CCT CAC AGG CGT TTG TAT GAT GTT GAC GAT GAG AAG ACC GTC CTG Asp Pro His Arg Arg Leu Tyr Asp Val Asp Asp Glu Lys Thr Val Leu 150 160	934
ATC ATC GGT GAC TGG TAT CAT GAA TCG TCC AAG GCA ATC CTT GCT TCT Ile Ile Gly Asp Trp Tyr His Glu Ser Ser Lys Ala Ile Leu Ala Ser 165 170 175 180	982
GGT AAC ATT ACC CGA CAG GTAAGTGATA CATGCCGGTC CCAGAAAAAT Gly Asn Ile Thr Arg Gln 185	1030
TCTCTAAATT CATTTAATT ACAG CGA CCG GTC TCT GCC ACC ATC AAC GGC Arg Pro Val Ser Ala Thr Ile Asn Gly 190 195	1081
AAA GGT CGA TTT GAC CCT GAC AAC ACT CCT GCC AAC CCA GAT ACT CTG Lys Gly Arg Phe Asp Pro Asp Asn Thr Pro Ala Asn Pro Asp Thr Leu 200 205 210	1129
TAC ACC CTC AAG GTC AAG CGA GGG AAG CGC TAT CGT CTG CGT GTC ATC Tyr Thr Leu Lys Val Lys Arg Gly Lys Arg Tyr Arg Leu Arg Val Ile 215 220 225	1177
AAT AGC TCG GAG ATC GCT TCG TTC CGA TTC AGT GTG GAA GGT CAC AAG Asn Ser Ser Glu Ile Ala Ser Phe Arg Phe Ser Val Glu Gly His Lys 230 235 240	1225
GTG ACT GTG ATT GCT GCC GAT GGC GTC TCT ACC AAA CCG TAT CAG GTC Val Thr Val Ile Ala Ala Asp Gly Val Ser Thr Lys Pro Tyr Gln Val 245 250 255	1273
GAT GCG TTT GAT ATT CTA GCA GGA CAG CGC ATA GAT TGC GTC Asp Ala Phe Asp Ile Leu Ala Gly Gln Arg Ile Asp Cys Val 260 265 270	1315
GTAAGTGTCG TCCGAACCCA CATCTGAGCT CAAGTGTTGA TACATGCGCG CTTATAG	1372
GTG GAG GCG AAC CAA GAA CCC GAC ACA TAC TGG ATC AAC GCA CCG CTG Val Glu Ala Asn Gln Glu Pro Asp Thr Tyr Trp Ile Asn Ala Pro Leu 275 280 285	1420
ACC AAC GTG CCC AAC AAG ACC GCT CAG GCT CTC CTC GTT TAT GAG GAG Thr Asn Val Pro Asn Lys Thr Ala Gln Ala Leu Leu Val Tyr Glu Glu 290 295 300 305	1468
GAT CGT CGG CCG TAC CAC CCT CCA AAG GGC CCG TAT CGC AAG TGG AGC Asp Arg Arg Pro Tyr His Pro Pro Lys Gly Pro Tyr Arg Lys Trp Ser 310 315 320	1516
GTC TCT GAG GCG ATC ATC AAG TAC TGG AAT CAC AAG CAC AAG CAC GGA Val Ser Glu Ala Ile Ile Lys Tyr Trp Asn His Lys His Lys His Gly 325 330 335	1564
CGT GGT TTG CTG TCT GGA CAT GGA GGT CTC AAG GCT CGG ATG ATC GAG Arg Gly Leu Leu Ser Gly His Gly Gly Leu Lys Ala Arg Met Ile Glu 340 345 350	1612
GGT AGC CAT CAT CTG CAT TCG CGC AGC GTC GTT AAG CGC CAG AAT GAG	1660

Gly Ser His His Leu His Ser Arg Ser Val Val Lys Arg Gln Asn Glu 355 360 365	
ACC ACC ACT GTT GTA ATG GAC GAG AGC AAG CTC GTT GTAAGTACCA Thr Thr Val Val Met Asp Glu Ser Lys Leu Val 370 375	1706
TATTTAAAAG TTGGTTGGGT TTCGAATACT TATTTCAACT TTTCTTAG CCA CTG GAA Pro Leu Glu	1763
TAC CCC GGC GCT GCA TGC GGG TCT AAA CCT GCT GAC CTC GTC TTG GAT Tyr Pro Gly Ala Ala Cys Gly Ser Lys Pro Ala Asp Leu Val Leu Asp 385 390 395 400	1811
CTC ACT TTT GGT TTG GTATGTAGCC AAATCGCCCA TATACAGGAT ACTGAATATT Leu Thr Phe Gly Leu 405	1866
GTTTGTGCGT GTAG AAC TTT GCT ACC GGG CAC TGG ATG ATC AAC GGT ATC Asn Phe Ala Thr Gly His Trp Met Ile Asn Gly Ile 410 415	1916
CCA TAC GAG TCT CCC AAA ATC CCC ACA TTG CTC AAG ATC CTC ACT GAT Pro Tyr Glu Ser Pro Lys Ile Pro Thr Leu Leu Lys Ile Leu Thr Asp 420 425 430	1964
GAG GAC GGG GTT ACC GAG TCT GAC TTC GTATGTTCCC TTTTCGGTAT Glu Asp Gly Val Thr Glu Ser Asp Phe 435 440	2011
CTTCGTATGC GTGCACTGAC TCGTGCTGGT GGGAATTTAG ACC AAG GAG GAG CAC Thr Lys Glu Glu His 445	2066
ACA GTC ATA CTC CCG AAG AAC AAA TGC ATC GAA TTC AAC ATC AAG GGG Thr Val Ile Leu Pro Lys Asn Lys Cys Ile Glu Phe Asn Ile Lys Gly 450 455 460	2114
AAC TCG GGT ATT CCC ATT ACG CAC CCC GTA CAT CTT CAC GGT Asn Ser Gly Ile Pro Ile Thr His Pro Val His Leu His Gly 465 470 475	2156
GTAAGTGCAT ATCGGATGGT TTACGATACT AAGGCTCATC AACTTTTTAG CAC ACT His Thr	2212
TGG GAT GTC GTA CAA TTT GGC AAC AAC CCA CCC AAT TAT GTC AAT CCT Trp Asp Val Val Gln Phe Gly Asn Asn Pro Pro Asn Tyr Val Asn Pro 480 485 490	2260
CCC CGT AGG GAC GTG GTT GGC TCT ACA GAT GCG GGT GTG AGG ATT CAG Pro Arg Arg Asp Val Val Gly Ser Thr Asp Ala Gly Val Arg Ile Gln 500 505	2308
TTC AAG ACC GAC AAT CCA GGA CCG TGG TTC CTG CAC TGC GTGCGTCGGT Phe Lys Thr Asp Asn Pro Gly Pro Trp Phe Leu His Cys 515 520	2357
CCCCATCGTC CGTTATGGTT TTTCTAATAC GTCCCATTCT ATTTTAG CAT ATT GAC His Ile Asp 525	2413
TGG CAT CTT GAG GAG GGT TTC GCA GTGAGTACTG AGACCTAAGT GCTACTCGGC Trp His Leu Glu Glu Gly Phe Ala	2467

PCT/US94/10264 WO 95/07988

TCATTACTGA TTACCGCATG TATGCGTCTA G ATG GTG TTT GCT GAA GCG CCC Met Val Phe Ala Glu Ala Pro 540	2519
GAA GCC GTC AAG GGA GGT CCA AAG AGC GTG GCC GTG GAC TCT CAG TGG Glu Ala Val Lys Gly Gly Pro Lys Ser Val Ala Val Asp Ser Gln Trp 545 550 555	2567
GAA GGG CTG TGT GGC AAG TAC GAC AAC TGG CTA AAA TCA AAT CCG GGC Glu Gly Leu Cys Gly Lys Tyr Asp Asn Trp Leu Lys Ser Asn Pro Gly 550 570	2615
CAG CTG TAGGCGTATC GCAGCCACAT TGGTGATGAT TGAAAGTTGC ATCTTGTTCC Gln Leu 575	2671
TATAACCGGC TCTTATATAC GGGTGTCTCC CAGTAAAGTC GTAGCCCAAT TTCAGCCGAG	2731
ACAGATATTT AGTGGACTCT TACTCTTGTG TCCCATTGAC GCACATCGTT GCATCAAACC	2791
TGCTTTTTAT CGTCCCTCTT TGTAATTTGT GTTGCTGTAA TGTATCG	2838

(2) INFORMATION FOR SEQ ID NO:2:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 576 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:2: Met Ala Arg Thr Thr Phe Leu Val Ser Val Ser Leu Phe Val Ser Ala Val Leu Ala Arg Thr Val Glu Tyr Gly Leu Lys Ile Ser Asp Gly Glu 20 25 30 Ile Ala Pro Asp Gly Val Lys Arg Asn Ala Thr Leu Val Asn Gly Gly Tyr Pro Gly Pro Leu Ile Phe Ala Asn Lys Gly Asp Thr Leu Lys Val Lys Val Gln Asn Lys Leu Thr Asn Pro Glu Met Tyr Arg Thr Thr Ser Ile His Trp His Gly Leu Leu Gln His Arg Asn Ala Asp Asp Asp Gly Pro Ser Phe Val Thr Gln Cys Pro Ile Val Pro Arg Glu Ser Tyr Thr

Tyr Thr Ile Pro Leu Asp Asp Gln Thr Gly Thr Tyr Trp Tyr His Ser 115 120 125

His Leu Ser Ser Gln Tyr Val Asp Gly Leu Arg Gly Pro Leu Val Ile

Tyr Asp Pro Lys Asp Pro His Arg Arg Leu Tyr Asp Val Asp Asp Glu

Lys Thr Val Leu Ile Ile Gly Asp Trp Tyr His Glu Ser Ser Lys Ala

PCT/US94/10264

.*

WO 95/07988 Ile Leu Ala Ser Gly Asn Ile Thr Arg Gln Arg Pro Val Ser Ala Thr 185 Ile Asn Gly Lys Gly Arg Phe Asp Pro Asp Asn Thr Pro Ala Asn Pro Asp Thr Leu Tyr Thr Leu Lys Val Lys Arg Gly Lys Arg Tyr Arg Leu Arg Val Ile Asn Ser Ser Glu Ile Ala Ser Phe Arg Phe Ser Val Glu Gly His Lys Val Thr Val Ile Ala Ala Asp Gly Val Ser Thr Lys Pro Tyr Gln Val Asp Ala Phe Asp Ile Leu Ala Gly Gln Arg Ile Asp Cys Val Val Glu Ala Asn Gln Glu Pro Asp Thr Tyr Trp Ile Asn Ala Pro Leu Thr Asn Val Pro Asn Lys Thr Ala Gln Ala Leu Leu Val Tyr Glu Glu Asp Arg Arg Pro Tyr His Pro Pro Lys Gly Pro Tyr Arg Lys Trp 305 310 315 Ser Val Ser Glu Ala Ile Ile Lys Tyr Trp Asn His Lys His Lys His Gly Arg Gly Leu Leu Ser Gly His Gly Gly Leu Lys Ala Arg Met Ile Glu Gly Ser His His Leu His Ser Arg Ser Val Val Lys Arg Gln Asn Glu Thr Thr Val Val Met Asp Glu Ser Lys Leu Val Pro Leu Glu Tyr Pro Gly Ala Ala Cys Gly Ser Lys Pro Ala Asp Leu Val Leu Asp 395 385 Leu Thr Phe Gly Leu Asn Phe Ala Thr Gly His Trp Met Ile Asn Gly Ile Pro Tyr Glu Ser Pro Lys Ile Pro Thr Leu Leu Lys Ile Leu Thr 425 Asp Glu Asp Gly Val Thr Glu Ser Asp Phe Thr Lys Glu Glu His Thr Val Ile Leu Pro Lys Asn Lys Cys Ile Glu Phe Asn Ile Lys Gly Asn Ser Gly Ile Pro Ile Thr His Pro Val His Leu His Gly His Thr Trp 475 470 Asp Val Val Gln Phe Gly Asn Asn Pro Pro Asn Tyr Val Asn Pro Pro 490 Arg Arg Asp Val Val Gly Ser Thr Asp Ala Gly Val Arg Ile Gln Phe Lys Thr Asp Asn Pro Gly Pro Trp Phe Leu His Cys His Ile Asp Trp 520

His Leu Glu Glu Gly Phe Ala Met Val Phe Ala Glu Ala Pro Glu Ala

530 535 540

Val Lys Gly Gly Pro Lys Ser Val Ala Val Asp Ser Gln Trp Glu Gly 545 550 555 560

Leu Cys Gly Lys Tyr Asp Asn Trp Leu Lys Ser Asn Pro Gly Gln Leu 565 570 575

- (2) INFORMATION FOR SEQ ID NO:3:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 3117 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: DNA (genomic)
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Rhizoctonia laccase
 - (ix) FEATURE:
 - (A) NAME/KEY: CDS
 - (B) LOCATION: join(393..524, 577..687, 737..799, 860..985, 1043 ..1045, 1097..1219, 1269..1538, 1601..1996, 2047 ..2118, 2174..2284, 2338..2439, 2495..2635, 2693 ..2725, 2786..2899)
 - (ix) FEATURE:
 - (A) NAME/KEY: intron
 - (B) LOCATION: 525..576
 - (ix) FEATURE:
 - (A) NAME/KEY: intron
 - (B) LOCATION: 688..736
 - (ix) FEATURE:
 - (A) NAME/KEY: intron
 - (B) LOCATION: 800..859
 - (ix) FEATURE:
 - (A) NAME/KEY: intron
 - (B) LOCATION: 986..1042
 - (ix) FEATURE:
 - (A) NAME/KEY: intron
 - (B) LOCATION: 1220..1268
 - (ix) FEATURE:
 - (A) NAME/KEY: intron
 - (B) LOCATION: 1539..1600
 - (ix) FEATURE:
 - (A) NAME/KEY: intron
 - (B) LOCATION: 1823..1936
 - (ix) FEATURE:
 - (A) NAME/KEY: intron
 - (B) LOCATION: 1973..2046
 - (ix) FEATURE:
 - (A) NAME/KEY: intron
 - (B) LOCATION: 2119..2173
 - (ix) FEATURE:
 - (A) NAME/KEY: intron

(B) LOCATION:	2285.	.2337
---------------	-------	-------

(ix) FEATURE:

(A) NAME/KEY: intron
(B) LOCATION: 2440..2494

(ix) FEATURE:

(A) NAME/KEY: intron
(B) LOCATION: 2636..2692

(iv) PEATURE:

(A) NAME/KEY: intron
(B) LOCATION: 1046..1096

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:	
GAGTGATCCG CCAGAGTTCA GGCGGATAAG TTCCTAAATA GTCATTCGCC TATTCGTGTA	60
CCTCAGCATA CTGACGACAT ACCGCCAGAT CGCCCTCGGT TCGGGCGTGG CATACGTTCG	120
CAAGGGCACC TCACGGAGCA AACTCTAAAA AGCTTCGGCA TGGATTGCAT TTTGTATTGT	180
AAACAAGTTA CGAGAAAAAC AATAGATCAG TTTTTGCCGA ATCGGATGGC TTGAAACGGA	240
AGTACCGATG GCCGATCCGA GTCGAATGAA TTAACGCATC TGAAACGGGA CCCTGAGTCG	300
AGGCACCCGC CGGCCTTGGC CGTATAAGTC ACTTGTCGCC AACTAGCACT TTTTCATTCC	360
CCCTTTTCTT CTTCCTCGTC TTCTTCTTCT CT ATG GCT CGG TCG ACT ACT TCA Met Ala Arg Ser Thr Thr Ser 1 5	413
CTC TTT GCA CTG TCT CTC GTT GCT TCA GCG TTT GCT CGA GTC GTT GAC Leu Phe Ala Leu Ser Leu Val Ala Ser Ala Phe Ala Arg Val Val Asp 10 15 20	461
TAT GGG TTT GAT GTG GCT AAT GGG GCA GTT GCT CCG GAT GGT GTA ACA Tyr Gly Phe Asp Val Ala Asn Gly Ala Val Ala Pro Asp Gly Val Thr 25 30 35	509
AGG AAC GCG GTT CTC GTGAGTTAGC TGTAAGATGG TGTATATGCT GGTTGCCTAA Arg Asn Ala Val Leu 40	564
CGGGAATGTC AG GTC AAT GGT CGC TTC CCT GGT CCA TTG ATC ACC GCC Val Asn Gly Arg Phe Pro Gly Pro Leu Ile Thr Ala 45 50 55	612
AAC AAG GGG GAT ACA CTT AAA ATC ACC GTG CGG AAT AAA CTC TCC GAT Asn Lys Gly Asp Thr Leu Lys Ile Thr Val Arg Asn Lys Leu Ser Asp 60 65 70	660
CCA ACT ATG CGA AGG AGC ACG ACC ATC GTTAGTACTT CCCCTCATCT Pro Thr Met Arg Arg Ser Thr Thr Ile 75 80	707
GTCTTGAAAC TTTCTCATCT TTTTTGAAG CAC TGG CAC GGT CTG CTC CAA CAC His Trp His Gly Leu Leu Gln His 85	760
AGG ACG GCA GAA GAA GAT GGC CCG GCC TTT GTA ACC CAG GTATGCCTTA Arg Thr Ala Glu Glu Asp Gly Pro Ala Phe Val Thr Gln 90 95 100	809
TCCTATCGCT GCTCTGTCCC CGCGTCCTTC CCTGACTCGG GCGATTCTAG TGC CCG Cys Pro	865

ATT CCT CCG CAA GAA TCG TAC ACC TAT ACG ATG CCG CTC GGC GAA CAG Ile Pro Pro Gln Glu Ser Tyr Thr Tyr Thr Met Pro Leu Gly Glu Gln 105 110 115 120	913
ACC GGC ACG TAT TGG TAC CAC AGC CAC TTG AGC TCC CAG TAT GTG GAC Thr Gly Thr Tyr Trp Tyr His Ser His Leu Ser Ser Gln Tyr Val Asp 125	961
GGG TTG CCT GGG CCC ATC GTT ATT GTAAGTCTTC ATTTAACCTT ATTCTTGGTT Gly Leu Arg Gly Pro Ile Val Ile 140	1015
ATGGCTGATT GTGACGTCGT GGTTAGT ATG TTCGTGGCTT CCACAAGAAG Met 145	1065
TCAGCAGCCC TTGAAGCTAA CTTTATTCCA G GAC CCC CAC GAC CCG TAC AGA Asp Pro His Asp Pro Tyr Arg 150	1117
AAC TAC TAT GAT GTC GAC GAC GAG CGT ACG GTC TTT ACT TTA GCA GAC Asn Tyr Tyr Asp Val Asp Asp Glu Arg Thr Val Phe Thr Leu Ala Asp 155 160 165	1165
TGG TAC CAC ACG CCG TCG GAG GCT ATC ATT GCC ACC CAC GAT GTC TTG Trp Tyr His Thr Pro Ser Glu Ala Ile Ile Ala Thr His Asp Val Leu 170 175 180	1213
AAA ACG GTACGCGTTA ATCCTTCTAG CTTTCTTTCC TTGGGTCACT TTCTATCAG Lys Thr 185	1268
ATC CCC GAC TCG GGT ACG ATC AAC GGC AAA GGC AAA TAC GAT CCT GCT Ile Pro Asp Ser Gly Thr Ile Asn Gly Lys Gly Lys Tyr Asp Pro Ala 190 195 200	1316
TCG GCT AAC ACC AAC AAC ACG ACA CTC GAG AAC CTC TAC ACT CTC AAA Ser Ala Asn Thr Asn Asn Thr Thr Leu Glu Asn Leu Tyr Thr Leu Lys 205 210 215	1364
GTC AAA CGC GGC AAG CGG TAT CGC CTG AGG ATT ATC AAC GCC TCG GCC Val Lys Arg Gly Lys Arg Tyr Arg Leu Arg Ile Ile Asn Ala Ser Ala 220 225 230	1412
ATC GCT TCG TTC CGG TTC GGC GTG CAG GGC CAC AAG TGC ACG ATC ATC Ile Ala Ser Phe Arg Phe Gly Val Gln Gly His Lys Cys Thr Ile Ile 235 240 245	1460
GAG GCT GAT GGC GTC CTC ACC AAA CCG ATC GAG GTC GAT GCG TTT GAT Glu Ala Asp Gly Val Leu Thr Lys Pro Ile Glu Val Asp Ala Phe Asp 255 260 265	1508
ATT CTA GCA GGC CAG AGG TAT AGC TGC ATC GTAAGTCTAC CTATGCCTTG Ile Leu Ala Gly Gln Arg Tyr Ser Cys Ile 270 275	1558
TTGTGGAGAT AAGAACCTGA CTGAATGTAT GCGCTCCAAT AG TTG AAG GCC GAC Leu Lys Ala Asp 280	1612
CAA GAT CCT GAT TCC TAC TGG ATA AAT GCG CCA ATC ACA AAC GTT CTC Gln Asp Pro Asp Ser Tyr Trp Ile Asn Ala Pro Ile Thr Asn Val Leu 285 290 295	1660
AAC ACC AAC GTC CAG GCA TTG CTA GTG TAT GAA GAT GAC AAG CGT CCT	1708

Asn Thr Asn Val Gln Ala Leu Leu Val Tyr Glu Asp Asp Lys Arg P 300 305 310	ro
ACT CAC TAC CCC TGG AAG CCG TTT TTG ACA TGG AAG ATA TCA AAT G Thr His Tyr Pro Trp Lys Pro Phe Leu Thr Trp Lys Ile Ser Asn G 315 320 325	
ATC ATT CAG TAC TGG CAG CAC AAG CAC GGG TCG CAC GGT CAC AAG G Ile Ile Gln Tyr Trp Gln His Lys His Gly Ser His Gly His Lys G 330 335 340	
AAG GGG CAT CAT CAT AAA GTC CGG GCC ATT GGA GGT GTA TCC GGG T Lys Gly His His Lys Val Arg Ala Ile Gly Gly Val Ser Gly L 345 350 355	TG 1852 Leu 160
AGC TCC AGG GTT AAG AGC CGG GCG AGT GAC CTA TCG AAG AAG GCT G Ser Ser Arg Val Lys Ser Arg Ala Ser Asp Leu Ser Lys Lys Ala V 365 370 375	FTC 1900 Val
GAG TTG GCT GCA CTC GTT GCG GGT GAG GCC GAG TTG GAC AAG A Glu Leu Ala Ala Leu Val Ala Gly Glu Ala Glu Leu Asp Lys A 380 385 390	
CAG AAT GAG GAT AAT TCG ACT ATT GTA TTG GAT GAG ACC AAG CTT A Gln Asn Glu Asp Asn Ser Thr Ile Val Leu Asp Glu Thr Lys Leu I 395 400 405	NTT 1996
GTAAGTCCCT TAATTTTTTT CGGTGTCACG GAAGCTAACC CGCGTAATAG CCG TT Pro Le 41	eu
GTT CAA CCT GGT GCA CCG GGC GGC TCC AGA CCA GCT GAC GTC GTG GVal Gln Pro Gly Ala Pro Gly Gly Ser Arg Pro Ala Asp Val Val V	
415 420 425	
415 420 425 CCT CTG GAC TTT GGC CTC GTATGTGGCT TCTTGTTATT CGTCCGGAAT Pro Leu Asp Phe Gly Leu 430	2148
CCT CTG GAC TTT GGC CTC GTATGTGGCT TCTTGTTATT CGTCCGGAAT Pro Leu Asp Phe Gly Leu	2148 ATA 2200
CCT CTG GAC TTT GGC CTC GTATGTGGCT TCTTGTTATT CGTCCGGAAT Pro Leu Asp Phe Gly Leu 430 GCAAACTGAT TTGGGTGGGC TATAG AAC TTT GCC AAC GGA CTG TGG ACG A Asn Phe Ala Asn Gly Leu Trp Thr I	2148 ATA 2200 Ele
CCT CTG GAC TTT GGC CTC GTATGTGGCT TCTTGTTATT CGTCCGGAAT Pro Leu Asp Phe Gly Leu 430 GCAAACTGAT TTGGGTGGGC TATAG AAC TTT GCC AAC GGA CTG TGG ACG A Asn Phe Ala Asn Gly Leu Trp Thr I 435 AAC AAT GTC TCC TAC TCC CCT CCG GAT GTC CCT ACT CTC CTC AAG A Asn Asn Val Ser Tyr Ser Pro Pro Asp Val Pro Thr Leu Leu Lys I	2148 ATA 2200 Ele
CCT CTG GAC TTT GGC CTC GTATGTGGCT TCTTGTTATT CGTCCGGAAT Pro Leu Asp Phe Gly Leu 430 GCAAACTGAT TTGGGTGGGC TATAG AAC TTT GCC AAC GGA CTG TGG ACG A Asn Phe Ala Asn Gly Leu Trp Thr I 435 AAC AAT GTC TCC TAC TCC CCT CCG GAT GTC CCT ACT CTC CTC AAG A Asn Asn Val Ser Tyr Ser Pro Pro Asp Val Pro Thr Leu Leu Lys I 445 TTG ACC GAC AAA GAC AAA GTC GAC GCT TCT GAC TTC GTAGGTTCCT Leu Thr Asp Lys Asp Lys Val Asp Ala Ser Asp Phe	2148 ATA 2200 Cle 2248 Cle 2294 GAA 2349
CCT CTG GAC TTT GGC CTC GTATGTGGCT TCTTGTTATT CGTCCGGAAT Pro Leu Asp Phe Gly Leu 430 GCAAACTGAT TTGGGTGGGC TATAG AAC TTT GCC AAC GGA CTG TGG ACG AAN Phe Ala Asn Gly Leu Trp Thr I 435 AAC AAT GTC TCC TAC TCC CCT CCG GAT GTC CCT ACT CTC CTC AAG AAN ASN Val Ser Tyr Ser Pro Pro Asp Val Pro Thr Leu Leu Lys I 445 TTG ACC GAC AAA GAC AAA GTC GAC GCT TCT GAC TTC GTAGGTTCCT Leu Thr Asp Lys Asp Lys Val Asp Ala Ser Asp Phe 460 CTTCTTCTTT TCAAACTAGC TACTGACATT AAGTGAACGT CAG ACG GCC GAT GTA AAC AAC GCC TACTGACATT AAGTGAACGT CAG ACG GCC GAT GTA AAC AAC AAC GCC TACTGACATT AAGTGAACGT CAG ACG GCC GAT GTACTGACATT AAGTGAACGT CAG ACG GCC GAT GTACTGA	2148 ATA 2200 The 2248 The 2294 SAA 2349 Shu AAG 2397
CCT CTG GAC TTT GGC CTC GTATGTGGCT TCTTGTTATT CGTCCGGAAT Pro Leu Asp Phe Gly Leu 430 GCAAACTGAT TTGGGTGGGC TATAG AAC TTT GCC AAC GGA CTG TGG ACG A Asn Phe Ala Asn Gly Leu Trp Thr I 435 AAC AAT GTC TCC TAC TCC CCT CCG GAT GTC CCT ACT CTC CTC AAG A Asn Asn Val Ser Tyr Ser Pro Pro Asp Val Pro Thr Leu Leu Lys I 445 TTG ACC GAC AAA GAC AAA GTC GAC GCT TCT GAC TTC GTAGGTTCCT Leu Thr Asp Lys Asp Lys Val Asp Ala Ser Asp Phe 460 CTTCTTCTTT TCAAACTAGC TACTGACATT AAGTGAACGT CAG ACG GCC GAT G Thr Ala Asp G CAC ACG TAT ATT CTT CCA AAG AAC CAA GTT GTC GAG TTG CAC ATC A His Thr Tyr Ile Leu Pro Lys Asn Gln Val Val Glu Leu His Ile I	2148 ATA 2200 The 2248 The 2294 SAA 2349 Shu AAG 2397

PCT/US94/10264 WO 95/07988

GCG Ala 505	TTC Phe	GAC Asp	GTC Val	GTC Val	CAA Gln 510	TTC Phe	GGC Gly	GAC Asp	AAC Asn	GCT Ala 515	CCA Pro	AAC Asn	TAC Tyr	GTG Val	AAC Asn 520	2545
CCT Pro	CCG Pro	CGT Arg	AGG Arg	GAT Asp 525	GTA Val	GTA Val	GGC	GTA Val	ACT Thr 530	GAT Asp	GCT Ala	GGA Gly	GTC Val	CGT Arg 535	ATC Ile	2593
CAG Gln	TTC Phe	AGA Arg	ACC Thr 540	GAT Asp	AAC Asn	CCG Pro	GGC Gly	CCT Pro 545	TGG Trp	TTC Phe	CTC Leu	CAT His	TGC Cys 55û			2635
GTAT	CTC	CTT (CATC	rccc2	AC CO	CTT	TTC	r TT	ACTT	ATGG	TTT	ACCT!	rgc (GATT'	rag	2692
CAC His	ATT Ile	GAT Asp	TGG Trp	CAC His 555	TTG Leu	GAA Glu	GAA Glu	GGA Gly	TTT Phe 560	GCT Ala	GTA	AGTT	ATT :	ATTC(CTATTC	2745
CGAJ	AGCA!	rcg (GGGA	GATG	CT A	ACCA	AGGG'	r gT(STTT.	raag	ATG Met	GTA Val	TTC Phe	GCC Ala 565	GAA Glu	2800
GCG Ala	CCT Pro	GAA Glu	GAT Asp 570	ATC Ile	AAG Lys	AAA Lys	GGC Gly	TCT Ser 575	CAG Gln	AGT Ser	GTC Val	AAG Lys	CCT Pro 580	GAC Asp	GGA Gly	2848
CAA Gln	TGG Trp	AAG Lys 585	Lys	CTA Leu	TGC Cys	GAG Glu	AAG Lys 590	Tyr	GAG Glu	AAG Lys	TTG Leu	CCT Pro 595	GAA Glu	GCA Ala	CTG Leu	2896
CAG Gln	TGA	AGTT	GCA	GTTG	TTTC	CC A	TTCG	GGAA	C TG	GCTC	ACTA	TTC	CTTT	TGC		2949
ATA	ATTC	GGA	CTTT	TATT	TT G	GGAC	ATTA	T TG	GACT	atgg	ACT	TGTT	TGT	CACA	CCCTCG	3009
CTC	ACTG	TGT	CCCT	CCTT	ga g	TACC	TATA	C TC	TATT	CGTA	TAG	TGGG	AAT	ATGG	AATATC	3069
GGA	TGTA	ATA	AATG	CTCG	TG C	CTTT	GGTG	C TC	GAAA	TGGG	GTA	GGAC	T			3117

(2) INFORMATION FOR SEQ ID NO:4:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 599 amino acids
 (B) TYPE: amino acid

 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:

Met Ala Arg Ser Thr Thr Ser Leu Phe Ala Leu Ser Leu Val Ala Ser

Ala Phe Ala Arg Val Val Asp Tyr Gly Phe Asp Val Ala Asn Gly Ala 20 25 30

Val Ala Pro Asp Gly Val Thr Arg Asn Ala Val Leu Val Asn Gly Arg

Phe Pro Gly Pro Leu Ile Thr Ala Asn Lys Gly Asp Thr Leu Lys Ile

Thr Val Arg Asn Lys Leu Ser Asp Pro Thr Met Arg Arg Ser Thr Thr 70

Ile His Trp His Gly Leu Leu Gln His Arg Thr Ala Glu Glu Asp Gly 85 90 95 Pro Ala Phe Val Thr Gln Cys Pro Ile Pro Pro Gln Glu Ser Tyr Thr Tyr Thr Met Pro Leu Gly Glu Gln Thr Gly Thr Tyr Trp Tyr His Ser His Leu Ser Ser Gln Tyr Val Asp Gly Leu Arg Gly Pro Ile Val Ile 130 140 Met Asp Pro His Asp Pro Tyr Arg Asn Tyr Tyr Asp Val Asp Asp Glu
145 150 155 160 Arg Thr Val Phe Thr Leu Ala Asp Trp Tyr His Thr Pro Ser Glu Ala Ile Ile Ala Thr His Asp Val Leu Lys Thr Ile Pro Asp Ser Gly Thr Ile Asn Gly Lys Gly Lys Tyr Asp Pro Ala Ser Ala Asn Thr Asn Asn Thr Thr Leu Glu Asn Leu Tyr Thr Leu Lys Val Lys Arg Gly Lys Arg 210 215 220 Tyr Arg Leu Arg Ile Ile Asn Ala Ser Ala Ile Ala Ser Phe Arg Phe Gly Val Gln Gly His Lys Cys Thr Ile Ile Glu Ala Asp Gly Val Leu Thr Lys Pro Ile Glu Val Asp Ala Phe Asp Ile Leu Ala Gly Gln Arg Tyr Ser Cys Ile Leu Lys Ala Asp Gln Asp Pro Asp Ser Tyr Trp Ile 275 280 285 Asn Ala Pro Ile Thr Asn Val Leu Asn Thr Asn Val Gln Ala Leu Leu 295 Val Tyr Glu Asp Asp Lys Arg Pro Thr His Tyr Pro Trp Lys Pro Phe 305 310 315 Leu Thr Trp Lys Ile Ser Asn Glu Ile Ile Gln Tyr Trp Gln His Lys His Gly Ser His Gly His Lys Gly Lys Gly His His Lys Val Arg Ala Ile Gly Gly Val Ser Gly Leu Ser Ser Arg Val Lys Ser Arg Ala Ser Asp Leu Ser Lys Lys Ala Val Glu Leu Ala Ala Ala Leu Val Ala Gly Glu Ala Glu Leu Asp Lys Arg Gln Asn Glu Asp Asn Ser Thr Ile 395 Val Leu Asp Glu Thr Lys Leu Ile Pro Leu Val Gln Pro Gly Ala Pro Gly Gly Ser Arg Pro Ala Asp Val Val Val Pro Leu Asp Phe Gly Leu 425 Asn Phe Ala Asn Gly Leu Trp Thr Ile Asn Asn Val Ser Tyr Ser Pro

445

5 · 440

Pro Asp Val Pro Thr Leu Leu Lys Ile Leu Thr Asp Lys Asp Lys Val 450 455

Asp Ala Ser Asp Phe Thr Ala Asp Glu His Thr Tyr Ile Leu Pro Lys 465 470 475

Asn Gln Val Val Glu Leu His Ile Lys Gly Gln Ala Leu Gly Ile Val 485 490 495

His Pro Leu His Leu His Gly His Ala Phe Asp Val Val Gln Phe Gly 500 505 510

Asp Asn Ala Pro Asn Tyr Val Asn Pro Pro Arg Arg Asp Val Val Gly 515 520 525

Val Thr Asp Ala Gly Val Arg Ile Gln Phe Arg Thr Asp Asn Pro Gly 530 540

Pro Trp Phe Leu His Cys His Ile Asp Trp His Leu Glu Glu Gly Phe 545 550 555 560

Ala Met Val Phe Ala Glu Ala Pro Glu Asp Ile Lys Lys Gly Ser Gln 565 570 575

Ser Val Lys Pro Asp Gly Gln Trp Lys Lys Leu Cys Glu Lys Tyr Glu 580 585 590

Lys Leu Pro Glu Ala Leu Gln 595

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:

Ala Val Arg Asn Tyr Lys Phe Asp Ile Lys Asn Val Asn Val Ala Pro 1 5 10 15

Asp Gly Phe Gln Arg Pro Ile Val Ser Val 20 25

- (2) INFORMATION FOR SEQ ID NO:6:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 20 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:

Ser Gln Tyr Val Asp Gly Leu Arg Gly Pro Leu Val Ile Tyr Asp Pro 1 10 15

Asp Asp Asp His 20

- (2) INFORMATION FOR SEQ ID NO:7:
 - (i) SEQUENCE CHARACTERISTICS:
 (A) LENGTH: 26 amino acids

- (B) TYPE: amino acid
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: peptide
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:7:

Ser Arg Tyr Asx Val Asx Asx Ala Ser Thr Val Val Met Leu Glu Asx

Trp Tyr Arg Thr Pro Ala Xaa Val Leu Glu

- (2) INFORMATION FOR SEQ ID NO:8:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 25 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:

Ser Leu Gly Pro Thr Pro Asn Tyr Val Asn Pro Xaa Ile Arg Asp Val

Val Arg Val Gly Gly Thr Thr Val Val

- (2) INFORMATION FOR SEQ ID NO:9:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 25 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:9:

Gly Leu Ala Leu Val Phe Ala Glu Ala Pro Ser Gln Ile Arg Gln Gly

Val Gln Ser Val Gln Pro Asp Asp Ala

- (2) INFORMATION FOR SEQ ID NO:10:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 14 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: peptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:10:

Ile Arg Tyr Val Gly Gly Pro Ala Val Xaa Arg Ser Val Ile 1 5 10

- (2) INFORMATION FOR SEQ ID NO:11:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 6 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide

	(xi)	SEQ	UENC	E DE	SCRI	PTIO	N: S	EQ I	D NO	:11:						
	Ile 1	Leu	Ala	Asn	Pro 5	Ala										
(2)	INFO	RMAT	ION	FOR	SEQ	ÌD N	0:12	:								
	(i)	(A (B (C	UENC) LE) TY) ST	ngth Pe: Pand	: 8 amin EDNE	amin o ac SS:	o ac id sing	ids								
	(ii)	MOL	ECUL	E TY	PE:	pept	ide									
	(xi)	SEQ	UENC	E DE	SCRI	PTIC	N: S	EQ I	D NO	:12:						
	Tyr 1	Glu	Ala	Pro	Ser 5	Leu	Pro	Thr	•							
(2)	INFO	RMAI	'ION	FOR	SEQ	ID N	10:13	:								
	(i)	(Ā (E (C	UENC () LE () TY () SI () TO	NGTH PE: RAND	: 19 nucl EDNE	12 t eic SS:	ase ació sing	pair l	s							
	(ii)	MOI	ECUL	E TY	PE:	cDNA										
	(vi)	ORI	GINA () OF	L SC GANI	URCE SM:	: Rhiz	octo	nia	laco	ase						
	(xi)	(2	ATURE A) NA B) LC	ME/K	EY:	CDS 85	1671	L								
	(xi)	SEÇ	UENC	E DE	ESCRI	PTIC	N: 5	SEQ 1	D NO	:13:	:					
CTA	ACGCI	TG	TGCC	GAGC	T CO	GATO	CAC	r agi	PAACO	CGC	GCC	\GTG1	GC 1	rggai	ATTCGC	60
GGC	CGCGI	CG 1	CACC	CTCCI	T C		ATG (Met I 1									111
TTG Leu 10	CTC Leu	GCT Ala	GCG Ala	GTC Val	TCA Ser 15	ACC Thr	CCC Pro	GCC Ala	TTT Phe	GCT Ala 20	GCC Ala	GTC Val	CGC Arg	AAC Asn	TAT Tyr 25	159
AAG Lys	TTC Phe	GAC Asp	ATC Ile	AAG Lys 30	AAC Asn	GTC Val	AAT Asn	GTC Val	GCT Ala 35	CCC Pro	GAT Asp	GGC Gly	TTT Phe	CAG Gln 40	CGC Arg	207
TCT Ser	ATC Ile	GTC Val	TCC Ser 45	GTC Val	AAC Asn	GGT Gly	TTA Leu	GTT Val 50	CCT Pro	GGC Gly	ACG Thr	TTG Leu	ATC Ile 55	ACG Thr	GCC Ala	255
AAC Asn	AAG Lys	GGT Gly 60	GAC Asp	ACC Thr	TTG Leu	CGC Arg	ATT Ile 65	AAT Asn	GTC Val	ACG Thr	AAT Asn	CAA Gln 70	CTC Leu	ACG Thr	GAC Asp	303
CCT Pro	AGT Ser 75	ATG Met	CGT Arg	CGT Arg	GCC Ala	ACA Thr 80	ACG Thr	ATT Ile	CAT His	TGG Trp	CAT His 85	GGA Gly	TTG Leu	TTC Phe	CAA Gln	351

GCT Ala 90	ACT Thr	ACC Thr	GCC Ala	GAC Asp	GAG Glu 95	GAT Asp	GGC Gly	CCC Pro	GCA Ala	TTC Phe 100	GTC Val	ACG Thr	CAA Gln	TGC Cys	CCT Pro 105	399
ATT Ile	GCG Ala	CAA Gln	AAT Asn	TTG Leu 110	TCC Ser	TAT Tyr	ACA Thr	TAC Tyr	GAG Glu 115	ATC Ile	CCA Pro	TTG Leu	CGC Arg	GGC Gly 120	CAA Gln	447
ACA Thr	GGA Gly	ACC Thr	ATG Met 125	TGG Trp	TAT Tyr	CAC His	GCC Ala	CAT His 130	CTT Leu	GCG Ala	AGT Ser	CAA Gln	TAT Tyr 135	GTC Val	GAT Asp	495
GGA Gly	TTG Leu	CGA Arg 140	GGC Gly	CCT Pro	TTG Leu	GTC Val	ATC Ile 145	TAT Tyr	GAT Asp	CCA Pro	AAC Asn	GAC Asp 150	CCA Pro	CAC His	AAG Lys	543
TCG Ser	CGC Arg 155	TAC Tyr	GAC Asp	GTG Val	GAT Asp	GAT Asp 160	GCG Ala	AGC Ser	ACA Thr	GTA Val	GTC Val 165	ATG Met	CTT Leu	GAG Glu	GAC Asp	591
TGG Trp 170	TAC Tyr	CAT His	ACT Thr	CCG Pro	GCA Ala 175	CCC Pro	GTT Val	CTA Leu	GAA Glu	AAG Lys 180	CAA Gln	ATG Met	TTC Phe	TCG Ser	ACT Thr 185	639
AAT Asn	AAC Asn	ACC Thr	GCT Ala	CTG Leu 190	CTC Leu	TCT Ser	CCT Pro	GTT Val	CCG Pro 195	GAC Asp	TCG Ser	GGT Gly	CTT Leu	ATC Ile 200	AAT Asn	687
GGC Gly	AAA Lys	GGG Gly	CGC Arg 205	TAT Tyr	GTG Val	GGC Gly	GGT Gly	CCC Pro 210	GCA Ala	GTT Val	CCC Pro	CGG Arg	TCA Ser 215	GTA Val	ATC Ile	735
AAC Asn	GTA Val	AAA Lys 220	Arg	GGG Gly	YYY Lys	CGA Arg	TAT Tyr 225	CGC Arg	TTG Leu	CGC Arg	GTA Val	ATC Ile 230	AAC Asn	GCT Ala	TCT Ser	783
GCT Ala	ATC Ile 235	GGG	TCG Ser	TTT Phe	ACC Thr	TTT Phe 240	TCG Ser	ATC Ile	GAA Glu	GGA Gly	CAT His 245	AGT Ser	CTG Leu	ACT Thr	GTC Val	831
Ile 250	Glu	Ala	Asp	Gly	ATC Ile 255	Leu	His	Gln	Pro	Leu 260	Ala	Val	Asp	Ser	Phe 265	879
Gln	Ile	Tyr	Ala	Gly 270		Arg	Tyr	Ser	Val 275	Ile	Val	Glu	Ala	Asn 280	Gln	927
Thr	Ala	Ala	Asn 285	Tyr	TGG	Ile	Arg	Ala 290	Pro	Met	Thr	Val	Ala 295	Gly	Ala	975
Gly	Thr	300	Ala	Asn	TTG	Asp	9ro 305	Thr	Asn	Val	Phe	310	Val	Leu	His	1023
TAC	GAG Glu 315	Gly	A GCG Ala	CCC Pro	AAC Asn	GCC Ala 320	Glu	CCC Pro	ACG Thr	ACG Thr	GAA Glu 325	Gln	GGC Gly	AGT Ser	GCT Ala	1071
ATC 11e 330	: Gly	Thi	GCA Ala	CTC Lev	GTI Val 335	Glu	GAG Glu	AAC Asn	CTC Leu	CAT His 340	Ala	CTC Leu	ATC Ile	AAC Asn	Pro 345	1119
GG(Gly	GCT Ala	CCC Pro	G GGC G Gly	GG(Gly 35(, Ser	GCT Ala	Pro	GCA Ala	A GAC Asp 355	Va]	TCC Ser	CTC Lev	AAT Asr	CTI Leu 360	GCA Ala	1167

ATT Ile	GGG Gly	CGC Arg	AGC Ser 365	ACA Thr	GTT Val	GAT Asp	GGG Gly	ATT Ile 370	CTT Leu	AGG Arg	TTC Phe	ACA Thr	TTT Phe 375	AAT Asn	AAC Asn	1215
ATC Ile	AAG Lys	TAC Tyr 380	GAG Glu	GCT Ala	CCT Pro	TCG Ser	TTG Leu 385	CCC Pro	ACG Thr	CTC Leu	TTG Leu	AAG Lys 390	ATT Ile	TTG Leu	GCA Ala	1263
AAC Asn	AAT Asn 395	GCG Ala	AGC Ser	AAT Asn	GAC Asp	GCC Ala 400	GAT Asp	TTC Phe	ACG Thr	CCA Pro	AAT Asn 405	GAG Glu	CAC His	ACT Thr	ATC Ile	1311
GTA Val 410	TTG Leu	CCA Pro	CAC His	AAT Asn	AAA Lys 415	GTT Val	ATC Ile	GAG Glu	CTC Leu	AAT Asn 420	ATC Ile	ACC Thr	GGA Gly	GCT Gly	GCA Ala 425	1359
GAC Asp	CAC His	CCT Pro	ATC Ile	CAT His 430	CTC Leu	CAC His	GGC Gly	CAT His	GTG Val 435	TTT Phe	GAT Asp	ATC Ile	GTC Val	AAA Lys 440	TCA Ser	1407
CTC Leu	GGT Gly	GGT Gly	ACC Thr 445	CCG Pro	AAC Asn	TAT Tyr	GTC Val	AAC Asn 450	CCG Pro	CCA Pro	CGC Arg	AGG Arg	GAC Asp 455	GTA Val	GTT Val	1455
CGT Arg	GTC Val	GGA Gly 460	GGC Gly	ACC Thr	GGT Gly	GTG Val	GTA Val 465	CTC Leu	CGA Arg	TTC Phe	AAG Lys	ACC Thr 470	GAT Asp	AAC Asn	CCA Pro	1503
GGC Gly	CCA Pro 475	TGG Trp	TTT Phe	GTT Val	CAC His	TGC Cys 480	CAC His	ATT Ile	GAC Asp	TGG Trp	CAC His 485	TTG Leu	GAG Glu	GCT Ala	GGG Gly	1551
CTC Leu 490	GCA Ala	CTT Leu	GTC Val	TTT Phe	GCC Ala 495	GAG Glu	GCC Ala	CCC	AGC Ser	CAG Gln 500	ATT Ile	CGC Arg	CAG Gln	GGT Gly	GTC Val 505	1599
CAG Gln	TCG Ser	GTC Val	CAG Gln	CCC Pro 510	AAC Asn	AAT Asn	GCC Ala	TGG Trp	AAC Asn 515	CAG Gln	CTC Leu	TGC Cys	CCC	AAG Lys 520	TAC Tyr	1647
	GCT Ala			Pro												1672

(2) INFORMATION FOR SEQ ID NO:14:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 529 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:14:

Met Leu Ser Ser Ile Thr Leu Leu Pro Leu Leu Ala Ala Val Ser Thr 1 5 10 15

Pro Ala Phe Ala Ala Val Arg Asn Tyr Lys Phe Asp Ile Lys Asn Val 20 25 30

Asn Val Ala Pro Asp Gly Phe Gln Arg Ser Ile Val Ser Val Asn Gly 35 40 45

Leu Val Pro Gly Thr Leu Ile Thr Ala Asn Lys Gly Asp Thr Leu Arg 50 55 60

Ile 65	Asn	Val	Thr	Asn	Gln 70	Leu	Thr	Asp	Pro	Ser 75	Met	Arg	Arg	Ala	Thr 80
Thr	Ile	His	Trp	His 85	Gly	Leu	Phe	Gln	Ala 90	Thr	Thr	Ala	qzA	Glu 95	Asp
Gly	Pro	Ala	Phe 100	Val	Thr	Gln	Суз	Pro 105	Ile	Ala	Gln	Asn	Leu 110	Ser	Tyr
Thr	Tyr	Glu 115	Ile	Pro	Leu	Arg	Gly 120	Gln	Thr	Gly	Thr	Met 125	Trp	Tyr	His
Ala	His 130	Leu	Ala	Ser	Gln	Tyr 135	Val	Asp	Gly	Leu	Arg 140	Gly	Pro	Leu	Val
Ile 145	Tyr	Asp	Pro	Asn	Asp 150	Pro	His	Lys	Ser	Arg 155	Tyr	Asp	Val	Asp	Asp 160
Ala	Ser	Thr	Val	Val 165	Met	Leu	Glu	Asp	Trp 170	Tyr	His	Thr	Pro	Ala 175	Pro
Val	Leu	Glu	Lys 180	Gln	Met	Phe	Ser	Thr 185	Asn	Asn	Thr	Ala	Leu 190	Leu	Ser
Pro	Val	Pro 195	Asp	Ser	Gly	Leu	Ile 200	Asn	Gly	Lys	Gly	Arg 205	Tyr	Val	Gly
Gly	Pro 210	Ala	Val	Pro	Arg	Ser 215	Val	Ile	Asn	Val	Lys 220	Arg	Gly	Lys	Arg
Tyr 225	Arg	Leu	Arg	Val	Ile 230	Asn	Ala	Ser	Ala	Ile 235	Gly	Ser	Phe	Thr	Phe 240
Ser	Ile	Glu	Gly	His 245	Ser	Leu	Thr	Val	Ile 250	Glu	Ala	Asp	Gly	Ile 255	Leu
His	Gln	Pro	Leu 260	Ala	Val	Asp	Ser	Phe 265	Gln	Ile	Tyr	Ala	Gly 270	Gln	Arg
_		275			Glu		280					285			
	290			,	Val	295					300				
305					Ala 310					315					320
				325					330				•	335	
			340		Leu			345					350		
		355			Leu		360					365			
Ī	370				Thr	375					380				
385			•		1 Lys 390					395	•				400
_				405					410	+				415	
Ile	Glu	Leu	Asn	Ile	Thr	Gly	Gly	Ala	Asp	His	Pro	Ile	His	Leu	His

420 425 430

Gly His Val Phe Asp Ile Val Lys Ser Leu Gly Gly Thr Pro Asn Tyr 435 440 445

Val Asn Pro Pro Arg Arg Asp Val Val Arg Val Gly Gly Thr Gly Val 450 455 460

Val Leu Arg Phe Lys Thr Asp Asn Pro Gly Pro Trp Phe Val His Cys 465 470 475

His Ile Asp Trp His Leu Glu Ala Gly Leu Ala Leu Val Phe Ala Glu 485 490 495

Ala Pro Ser Gln Ile Arg Gln Gly Val Gln Ser Val Gln Pro Asn Asn 500 505 510

Ala Trp Asn Gln Leu Cys Pro Lys Tyr Ala Ala Leu Pro Pro Asp Leu 515 520 525

Gln

What we claim is:

- 1. A nucleic acid fragment containing a nucleic acid sequence encoding a Rhizoctonia laccase which functions optimally at pH between about 6.0 and 8.5.
 - 2. The fragment of Claim 1 which comprises a sequence encoding a Rhizoctonia solani laccase.
- 10 3. The fragment of Claim 1 which comprises a nucleic acid sequence encoding the amino acid sequence depicted in SEQ ID NO. 2.
- The fragment of Claim 1 which comprises a nucleic acid
 sequence encoding the amino acid sequence depicted in SEQ ID
 NO. 4.
- 5. The fragment of Claim 1, which comprises a nucleic acid sequence encoding a protein containing one or more of the amino acid sequences depicted in SEQ. ID NOS. 5, 6, 7, 8, 9, 10, 11, or 12.
- 6. The fragment of Claim 1 which comprises a nucleic acid sequence encoding the amino acid sequence depicted in SEQ ID 25 NO. 14.
 - 7. The fragment of Claim 1, which comprises the nucleic acid sequence depicted in SEQ ID NO. 1.
- 30 8. The fragment of Claim 1, which comprises the nucleic acid sequence depicted in SEQ. ID. NO. 3.

9. The fragment of Claim 1, which comprises the nucleic acid sequence depicted in SEQ. ID. NO. 13.

- 10. The fragment of Claim 1, which comprises the nucleic acid sequence contained in NRRL B-21141.
 - 11. The fragment of Claim 1, which comprises the nucleic acid sequence contained in NRRL B-21142.
- 10 12. The fragment of Claim 1, which comprises the nucleic acid sequence encoding the laccase produced by RS 22.
 - 13. The fragment of Claim 1, which comprises the nucleic acid sequence contained in NRRL B-21156.
- 14. A substantially pure *Rhizoctonia* laccase enzyme which functions optimally at a pH between about 6.0-8.5.

15

- 15. The enzyme of Claim 14 which is a Rhizoctonia solani 20 laccase.
 - 16. The enzyme of Claim 14 which comprises the sequence depicted in SEQ ID NO. 2, or a sequence with at least 80% homology thereto.
 - 17. The enzyme of Claim 14 which comprises the sequence depicted in SEQ ID NO 4, or a sequence with at least 80% homology thereto.
- 18. The enzyme of Claim 14 which comprises one or more of the peptide sequences depicted in SEQ ID NOS.5, 6, 7,

8, 9, 10, 11 or 12, or a sequence with at least 80% homology to one or more of these peptides.

- 19. The engyme of Claim 14 which comprises the sequence 5 depicted in SEQ ID NO 14, or a sequence with at least 80% homology thereto.
- 20. A recombinant vector comprising a nucleic acid fragment containing a nucleic acid sequence encoding a *Rhizoctonia*10 laccase which functions optimally at pH between about 6.0-8.5.
 - 21. The vector of Claim 20 in which the fragment is operably linked to a promoter sequence.

15

- 22. The vector of Claim 21 in which the promoter is a fungal or yeast promoter.
- 23. The vector of Claim 22 in which the promoter is the 20 TAKA amylase promoter of Aspergillus oryzae.
 - 24. The vector of Claim 22 in which the promoter is the glucoamylase (gluA) promoter of Aspergillus niger or Aspergillus awamsii.
 - 25. The vector of Claim 21 which also comprises a selectable marker.
- 26. The vector of Claim 25 in which the selectable marker is the amdS marker of Aspergillus nidulans or Aspergillus oryzae.

27. The vector of Claim 25 in which the selectable marker is the pyrG marker of Aspergillus nidulans, Aspergillus niger, Aspergillus awamorii, or Aspergillus oryzae.

- 5 28. The vector of Claim 21 which comprises both the TAKA amylase promoter of Aspergillus oryzae and the amdS or pyrG marker of Aspergillus nidulans or Aspergillus oryzae.
- 29. A host cell comprising a heterologous nucleic acid
 10 fragment containing a nucleic acid sequence encoding a
 Rhizoctonia laccase which functions optimally at pH between
 about 6.0-8.5.
 - 30. The host cell of Claim 28 which is a fungal cell.

15

- 31. The host cell of Claim 30 which is an Aspergillus cell.
- 32. The host cell of Claim 29 in which the fragment is integrated into the host cell genome.

- 33. The host cell of Claim 29 in which the fragment is contained on a vector.
- 34. The host cell of Claim 29 which comprises a fragment containing a sequence encoding the amino acid sequence depicted in SEQ ID NO. 2.
- 35. The host cell of Claim 29 which comprises a fragment containing a sequence encoding the amino acid sequence 30 depicted in SEQ ID NO: 4.

- 36. The host cell of Claim 29 which comprises a fragment containing a sequence encoding the amino acid sequence depicted in SEQ ID NO: 14.
- 5 37. The host cell of Claim 29 which comprises a fragment containing a sequence encoding one or more of the amino acid sequences depicted in SEQ ID NOS.: 5, 6, 7, 8, 9, 10, 11, or 12.
- 38. A method for obtaining a laccase enzyme which functions optimally at a pH between about 6.0-8.5 which comprises culturing a host cell comprising a nucleic acid fragment containing a nucleic acid sequence encoding a *Rhizoctonia* laccase enzyme which functions optimally at a pH between about 6.0-8.5, under conditions conducive to expression of the enzyme, and recovering the enzyme from the culture.
- 39. A method for polymerizing a lignin or lignosulfate substrate in solution which comprises contacting the substrate with a *Rhizoctonia* laccase which functions optimally at a pH between about 6.0-8.5.
- 40. A method for in situ depolymerization in Kraft pulp which comprises contacting the pulp with a *Rhizoctonia*25 laccase which functions optimally at a pH between about 6.0-8.5.
- 41. A method for oxidizing dyes which comprises contacting the dye with a *Rhizoctonia* laccase which functions optimally 30 at a pH between about 6.0-8.5.

42. A method of polymerizing a phenolic compounds which comprises contacting the phenolic compound with a *Rhizoctonia* laccase which functions optimally at a pH between about 6.0-8.5.

1 AGCGTCACACCAGACATCGGATGAAACGGAAAGGAAGGAA	GCAC 180 R T 4	AGTA 240 E Y 24	CTTT 300 T L 44	TGGA 360	CCAA 420 Q 67	tatc 480 81	AGAA 540 R 90
61 AACCACTGTTCATCTCGCGAGCTAACATGGGCGACGTATAAGAAGAACGCGAGAATGGGC	121 AGATTTCGATATCCCCTCTCGTCTCGGTTTTTGGTCTCGGCTTGCCTCTAATGGCGCGCAC	181 CACTITICETTICGETITICGETETTIGETITICGETGETGETA 4 T F L V S V S L F V S A V L A R T V E Y	241 CGGCTTGAAGATTAGTGATGGGAGATAGCTCCTGACGGTGTTAAGCGTAATGCGACTTT 24 G L K I S D G E I A P D G V K R N A T L	\sim N $_{301}$ GGgtacgcactccttgtaatccaacaattcaaggtttctgatgcttggtcagTAAATGGA \sim 44	361 GGGTATCCCGGTCCACTTTTTGCCAACAAGGGGGATACTCTCAAAGTCAAGGTCCAA 47 G Y P G P L I F A N K G D T L K V K V Q	421 AACAAGCTCACGAATCTGTATCGCACCACTTCCATCGtatgttcgttcgatatc	L tactaatacatccgtcgctaaatatcttgtagCATTGGCACGGTCTCTTACAACATAGAA
, ,	12.	18.	24 2 2	0° 4 / 21	36.	42.	481 81

F I G. 1A

1020 185

960 172

S

3

Ö

Н

H

h

×

田

Ω

Ω

>

Ω

901 152

TITGTATGATGATGAGAAGACCGTCCTGATCATCGGTGACTGGTATCATGAATC

GTCCAAGGCAATCCTTGCTTCTGGTAACATTACCCGACAgtaagtgatacatgccggtcc

Ø

ĸ

H

 $\boldsymbol{\mathsf{H}}$

Z

r

Ŋ

J

Ŋ

961

102	660	720	780 144	840 145	900
541 ACGCCGACGACGGTCCTTCGTCACTCAGGtaggattctggaaggttggcctga 90 N A D D G P S F V T Q	601 actctctgttaaccgacaacccgatgtcaccagTGCCCGATTGTTCCACGCGAGTCGTAT C P I V P R E S Y	661 ACTTACACCATACCTGGACGATCAAACCGGAACCTATTGGTACCATAGCCACTTGAGT 111 T Y T I P L D D Q T G T Y W Y H S H L S	721 TCGCAATACGTTGATGGTCTTCGAGGCCCGCTGGTAATCTgtgagtatcttgacttgtct 131 S Q Y V D G L R G P L V I	781 actgaaggcaacgagactaaaacaagcgtcgattcacagATGgttcgtctccctttatt 144	841 tagetetggatetteteteaegtaatacatgatagATCCCAAGGATCCTCACAGGCG
50	1(٦ و	7	7	7 00

1560 335

94	.40 214	1200 234	250 254	1320 272	1380 275	1440 295	1500 31.5
10	17		17	H	H		-
ဗ္ဗဗ	AA A	TT	AA X	lag	A GC	AC	ည်မှ
AAC N	CIC	ည်င် အ	ACC	gta	GAG	AAG	ည်
TC	T T	3CT A	ICT S	3TC V	GTG V	AAC	AAG K
T T	Y Y	I	STC V	ပ္တပ	:ag(P	ار الح
CCA	TGI	AGA E	ည်	ATT D	rat.	TGC V	CTC
CTG S	CTC	S S	ATG D	TAG	gct	ACG	ACC H
TCT	ATA D	GCT S	CCG	GCA	င်ရွင	CCA	ACC Y
CGG P	CAG P	ata N	CTG A	AGC Q	atg	TGA	CGT
ttcattttaattacagGCGACCGGTCTCTGCCATCAACGG 1030 R P V S A T I N G 194	CAAAGGTCGATTTGACCTGAACACTCCTGCCAACCCAGATACTCTGTACACCCTCAA 1140 K G R F D P D N T P A N P D T L Y T L K 214	GGTCAAGCGAAGCGCTATCGTCTGCGTCATCAATAGCTCGGAGATCGCTTCGTT V K R G K R Y R L R V I N S S E I A S F	CCGATTCAGTGTGAAGGTCACAAGGTGACTGTGATTGCTGCCGATGGCGTCTCTACCAA 1260 R F S V E G H K V T V I A A D G V S T K 254	ACCGTATCAGGTCGATGCGTTTGATATTCTAGCAGGACAGCGCATAGATTGCGTCGtaag	tgtcgtccgaacccacatctgagctcaagtgttgatacatgcgcgcttatagGTGGAGGC $_{ m V}$	GAACCAAGAACCCGACACATACTGGATCAACGCACCGCTGACAACAAGAC N Q E P D T Y W I N A P L T N V P N K T	CGCTCAGGCTCTCGTTTATGAGGAGGATCGTCGGCCGTACCACCCTCCAAAGGGCCCC
agc	A CCA	TCA <	TGA	SCAG A	tga	icac A	GTC R
aca	CTG	GTG	CTG	TAG	ıtgt	ACG	ATC D
att	CTC	.TGC	TGA V	I	aag	I	AGG E
tta	ACA	GTC	AGG K	ATE D	JCtC	M W	;AGG E
att	ACA	Y	AC? H	F	gaç	PACT Y	Y.
tto	CTG P	GCI	GTC G	, ₽	tct	CAJ T.	TTT V
cagaaaaattctctaaa	ACC D	AGC	AAG E	SATC D	ace	ACP D	J.
tct	'TTG F	GGA	TGG V	CGTATCAGGTCGA7 P Y Q V D	0001	P P	CTCAGGCTCTCCTC A Q A L L
tta	GAT R	GAG R	GTG S	AGG Q	gaa	AAC	SCTC A
ಇಇಇ	3TC G	AGC K	TCA	ATC	tac	AAG Q	AGG Q
jaa	A.A.G.	ICA.	3AT R	CGT P	tcg	ACC	CTC
cai	CAN	99		AC		GA	Ö
1021 185	1081 194	1141 214	1201 234	1261 254	5 1321 \$\sqrt{272}\$	1381 275	1441 295
ਜੋ``	₩.,	-	₩.	₽ .	3/2	. ← 1	7

S H 340	ATGGA 1680 M D 350	ttatt 1740	TGACC 1800 A D 365	stactg 1860 374	rcccar 1920 I P 387	rracce 1980 7 T 407	gctggt 2040 411	I E 427
	TCATCTGCATTCGCGCGCGTCGTTAAGCGCCAGAATGAGACCACCACTGTTGTAATGGA H L H S R S V V K R Q N E T T T V V M D		tcaacttttcttagCCACTGGAATACCCCGGCGCTGCATGCGGGTCTAAACCTGCTGACC	1801 TCGTCTTGGATCTCACTTTTGGTTTTGGtatgtagccaaatcgcccatatacaggatactg	aatattgtttgtgcgtgtagAACTTTGCTACCGGGCACTGGATGATCAACGGTATCCCAT N F A T G H W M I N G I P		1981 AGTCTGACTTgtatgttcccttttcggtatcttcgtatgcgtgcactgactcgtgctggt	2041 gggaatttagCACCAAGGAGGAGCACACAGTCATACTCCCGAAGAACAAATGCATCGAAT
1561 335	1621 340	1681 350	1741 350	1801 365	4 1861		1981 407	2041

F 1 G. 10

2:160	2220	2280	2340	2400	2460	2520	2580	2640
.146	451		491	493	504	511	531	545
TCAACATCAAGGGGAACTCGGGTATTCCCATTACGCACCCCGTACATCTTCACGGTGtaa F N I K G N S G I P I T H P V H L H G	gtgcatatcggatggtttacgatactaaggctcatcaactttttagCACACTTGGGATGT 2,320 H T W D V 451	CGTACAATTTGGCAACAACCCACCCAATTATGTCAATCCTCCCGTAGGGACGTGGTTGG V Q F G N N P P N Y V N P P R R D V V G	CTCTACAGATGCGGGTGTGAGGATTCAAGACCGACAATCCAGGACCGTGGTTCCT S T D A G V R I Q F K T D N P G P W F L	GCACTGgtgcgtcggtccgtcgftatggttttctaatacgtcccattctattt	tagCCATATTGACTGGCATCTTGAGGGGTTTCGCAAgtgagtactgagacctaagtgc H I D W H L E E G F A	tactcggctcattactgattaccgcatgtatgcgtctagTGGTGTTTGCTGAAGCGCCCG ${ t M}$	AAGCCGTCAAGGGTCCAAAGAGCGTGGCCGTGGACTCTCAGTGGGAAGGGCTGTGTG E A V K G G P K S V A V D S Q W E G L C	GCAAGTACGACAACTGGCTAAAATCAAATCCGGGCCAGCTGTAGGCGTATCGCAGCCACA G K Y D N W L K S N P G Q L *
2101	2161	2221	2281	2341	5 2401 4 93	2461	2521	2581
427	446	451	471	491		504	511	531

ATGATTGAAAGTTGCATCTTGTTCCTATAACCGGCTCTTATATATA	2700	2760	2820	2838	
2641 TTGGTG1 2701 CCAGTA1 2761 GTCCCA' 2821 TGTTGC'	2641 TIGGIGATGATIGAAAGTIGCATCTIGITCCIATAACCGGCTCTTATATACGGGTGTCTC	2701 CCAGTAAAGTCGTAGCCCAATTTCAGCCGAGACAGATATTTAGTGGACTCTTACTCTTGT	2761 GTCCCATTGACGCĂCATCGTTGCATCAAACCTGCTTTTTTATCGTCCCTCTTTTGTAATTTG	2821 TGTTGCTGTATCTATCG	

7	1 AAGCTTCGGCATGGATTGCATTTTGTATTGT	∩. 8 .T
181	181 AAACAAGTTACGAGAAAAAAATAGATCAGTTTTTGCCGAATCGGATGGCTTGAAACGGA	240
241	241 AGTACCGATGGCCGATCCGAGTCGAATTAACGCATCTGAAACGGGACCCTGAGTCG	300
301	301 AGGCACCCGCCGGCCTTGTAAGTCACTTGTCGCCAACTAGCACTTTTTCATTCC	360
361 1	361 CCCTTTTCTTCTTCTTCTTCTTCTTATGGCTCGACTACTTCACTCTTTG 1	420
421 10	421 CACTGTCTCTGGCCGCCCTTGGCTCGAGTCGTTGACTATGGGTTTGATGTGGCTA 10 A L S L A A P A L A R V V D Y G F D V A	480
481 30	481 ATGGGGCAGTTGCTCGGATGGTGTACAAGGAACGCGGTTCTCGgtgagttagctgtaa 30 N G A V A P D G V T R N A V L	540 45
541 45	gatggtgtatatgctggttgcctaacgggaatgtcagTCAATGGTCGCTTCCCTGGTCCA V N G R F P G P	600 53
601 53	601 TTGATCACCGCCAACAAGGGGGATACACTTTAAAATCACCGTGCGGAATAAACTCTCCGAT 53 L I T A N K G D T L K I T V R N K L S D	660

F 16.2A

CTCCGCAA P P Q ACCACAGC Y H S AGCACAGA P Y R ACCACACC	GAATCO E S CACTTO H L H L AACTAO N Y SCCGTCO	TTACACC Y T SAGCTCC SCTCT SCTATGAT Y D Y D SGAGGC'	TATAC Y T CAGTA Q Y CLEGG GCAGC V I I I	GATGCC M P TGTGGA Ctatgg ctttga CGACGA D E TGCCAC
CTCCGCAR P P Q ACCACAGC Y H S AGCACAGA P Y R ACCACACC	GAATCO ESCACTTV HL HL LEETAACAACAACAACAACAACAACAACAAACAAACAAACA	Y Y SAGC S S CCET Y Y Y Y	ACC T T TTCC S S SAT CAT D D	ctgactcgggcgattctagTGCCGATTCCTCGCAAGAATCGTACATACGATGCC 900 ctgactcgggcgattctagTGCCCGATTCCTCGCAAGAATCGTACACCTATACGATGCC 900 GTCGGCGAACAGACCGGCACTTTGTAGTACCACTTGAGCTCCCAGTATGTGGA 960 L G E Q T G T Y W Y H S H L S S Q Y V D 137 CGGGTTGCGTGGGCCCATCGTTATTTTGTAGAGCTCCCAGTATGTGGA 960 G L R G P I V I ctgattgtgacgtcgttggttagATGgttcgtggcttcacaaagaagtcagcagccttgg 1020 gctaactttattccagACCCCACGACCGTACAGAACTACTATGATGTCGACGACGA 1140 gcGTACGGTCTTTACTTTAGCAGACTGGTACCACGGGGCTATTGCCAC 1200 GCGTACGGTCTTTACTTTAGCAGACTGGTACCACGGCGTCGGAGGCTATTGCCAC 1200 R T V F T L A D W Y H T P S E A I I A T 180

F 16. 2B

 $\begin{array}{c} 1620 \\ 282 \end{array}$

Ω

gtggagataagaacctgactgaatgtatgcgctccaatag $_{
m L}$ K A D O D P

1561 275

TGATTCCTACTGGATAAATGCGCCAATCACAACGTTCTCAACACCCAACGTCCAGGCATT

1680 302

Z

Z

J.

Z

Д

Ø

Z

3

1621

282

1740 322

3

J

Д

3

H

H

Δ,

民

Ω

Ω

回

1681 302

1260 185	1320	1380	1440 242	1500 262	1560 275
1201 CCACGATGTCTTGAAAACgtacgcgttaatccttctagctttctttccttgggtcacttt 180 H D V L K T	1261 ctatcagGATCCCCGACTCGGGTACGATCAACGCCAAAAACGATCCTGCTTCGG 185 I P D S G T I N G K G K Y D P A S	1321 CTAACACCAACAACACTCGAGAACCTCTACACTCTCAAAGTCAAACGCGGCAAGC 202 A N T N N T T L E N L Y T L K V K R G K	1381 GGTATCGCCTGAGGATTATCAACGCCTCGGCCATCGCTTCGGTTCGGCGTGCAGG	1441 GCCACAAGTGCACGATCATCGAGGCTGATGGCGTCCTCACCAAACCGATCGAGGTCGATG 242 G H K C T I E A D G V L T K P I E V D	0 > 1501 CGTTTGATATTCTAGCAGGCCAGAGGTATAGCTGCATCGtaagtctacctatgccttgtt N 1501 CGTTTGATATTCTAGCAGGCCAGAGGTATAGCTGCATCG N 1501 CGTTTGATATTCTAGCAGGCCAGAGGTATAGCTGCATCG
1201 180	1261 185	1321 202	1381 222		5 7 1501 262
				•	- '

F I G. 2C

GCTAGTGTATGAAGATGACAAGCGTCCTACTCACTACCCCTGGAAGCCGTTTTTGACATG

342	1860 362	1920 349	1980 361	2040 361	2100 379	2160 335	2220 401	230
→				20			7	2,
A ×	AG R	GT V	SGA D	Jag	STC V	ה ה	ည်း	IGA D
H	17CC S	CIC	TT	ວີວວ	TGG V	gat	77C S	ည်း
ည် ပြွ	S S	3CA A	3TA V	aac		act	rac Y	GCT
ACC H	TG? L	CTC	TT1 I	Icta	DO C	สล	ည်	3AC D
CGC S	GGT	CTG A	CTA	aag	TGA	tgc	TCT V	TC(
3GT 3	SCG S		CGA S	299	AGC A	gaa	ATG N	AAG K
) -	TAT S	GT.	TTA	Ca	ACC.	Scgi	N N	ACA
(C)	1G1	CG2	TAT	ıtgt	AG7	gto	אאלי [AAG.
AX X	AGG G	rgr V	GGA D	G G G	ည်	tto	GAT	CA
S H H	0 0	3GC'	5 E	יו גר	ည်တ	t ta	3AC T	CGA
Q Q	AT	AAQ K	N N	יר דר	ည္တပ္သ	tgt)TG	3AC T
TG ≽	ეე ∀	AAG K	CAG	att	CGG P	tat	CTC	TT(L
rac Y	CGG R	ICG S	AGG R	tta	CAC	gct	GGA G	ATC
AG.	JIC(E L	AAG. K	SC	3TG	gtg	AAC	AAG K
TTI	GGGAAAGGGGCATCATAAAGTCCGGGCCATTGGAGGTGTATCCGGGTTGAGCTCCAG G K G H H H K V R A I G G V S G L S S R	GGTTAAGAGCCGGGCGAGTGACCTATCGAAGAAGGCTGTCGACTCGT V K S R A S D L S K K A V E L A A A L V	TGCGGGTGAGGCCGAGTAGAGGCAGAATGAGGATAATTCGACTATTGTATTGGA A G E A E L D K R Q N E D N S T I V L D	TGAGACCAAGCTTATT $oldsymbol{ t g}$ taatttttttt $oldsymbol{ t t}$ cog $oldsymbol{ t g}$ g $oldsymbol{ t g}$	taatagccgrrcgrcrcrcrcrcrcrcrcrcrcrcrcrcrcrcr	:atç	ŞÇÇ7	TCCGGATGTCCCTACTCTCAAGATCTTGACCGACAAAGACAAAGTCGACGCTTCTGA 2230 P D V P T L L K I L T D K D K V D A S D 421
TCA	ATA H	GTG S	ŢĞĞ L	taa	ACC	Cgt '	TTC)TC
AAA E	ATC	CGA	AGT E	TTg I	15 Q	CCI	ACT	T T
VTG	17C	3666	SCG	AGACCAAGCTTAT E T K L I	3GT V	r G	agA	CTA P
AAA	GC.	CCC) 1990 1	GC.	TI	F	ats	222
ATC	9 999	GAG	TGA	CAA	S P	GAC	gct	TG
BAT	AAA K	ľAA K	36G G	3AC T	tag	CTG	t gg	GGA
AAC K	GG2	GT.	CCC A	GA(aa	PCT	199 ⁽	ICC P
1741 GAAGATATCAATCATTCAGTACTGGCAGCACAGGGTCGCACGGTCACAA 1800 322 K I S N E I I Q Y W Q H K H G S H G H K 342				. H.	11 t	2101 CCTCTGGACTTTGGCCTCgtatgtggcttcttgttattcgtccggaatgcaaactgattt 379 P L D F G L	2161 gggtgggctatagAACTTTGCCAACGGACTGTGGACGATAAACAATGTCTCCTACTCCCC 385	221 401
1741 322	1801 342	1861 349	1921 349	1981 361	2 2041 361	210 37	216 38	2221 401
				10)/21			

F 16. 20

423	2400 453	2460 466	2520 475	2580 495	2640 513	2700 516	2760 524	2820 536	
	GCCGATGAACACGTATATTCTTCCAAAGAACCAAGTTGTCGAGTTGCACATCAAGGGA A D E H T Y I L P K N Q V V E L H I K G	CAGGCTTTGGGAATCGTACACCCCTTCATCTGCATGGCgtacgtcttctcacactgtt Q A L G I V H P L H L H G	ccagctcctattctctaacacactcctgcgatagCATGCGTTCGACGTCGTCCAATTCGG H A F D V V Q F G	CGACAACGCTCCAAACTACGTGAACCCTCCGCGTAGGGATGTAGTAGCGTAACTGATGC 2580 D N A P N Y V N P P R R D V V G V T D A 495	TGGAGTCCGTATCCAGTTCAGAACCGATAACCCGGGCCCTTGGTTCCTCCATTGGTALGC		TYGGCACTYGGAAGAATYYGCTAgtaagttattattcctattccgaagcatcgggga $forall W$ $forall H$ $forall E$ $forall E$ $forall A$	gatgctaaccaagggtgttttaagTGGTATTCGCCGAAGCGCCTGAAGATATCAAGAA 2820	
2281 421	23 <u>41</u> 423	2401 453	2461 466	2521 475	2/11 2581 11/2	2641 513	2701 516	2761 524	

2880 556	2940 562	3000	·	
2821 AGGCTCTCAGAGTGTCAAGCCTGACGACAATGGAAGAAACTATGCGAGAAGTATGAGAA 2880 536 G S Q S V K P D G Q W K K L C E K Y E K 556	2881 GTTGCCTGAAGTGAAGTTGCAGTTGTTTCCCATTCGGAACTGGCTCACTAT 556 L P E A L Q *	2941 TCCTTTTGCATAATTCGGACTTTTTTTGGGACATTATTGGACTATGCATTTGTTTTGTC 3000	3001 ACACCGCGGAACTAAGCCGAATTC	F I G. 2F

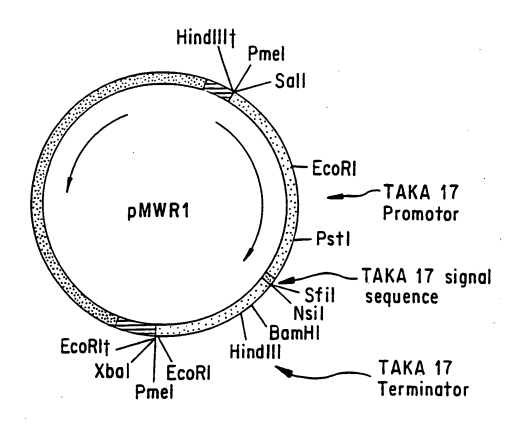


FIG. 3

132 GCC 	186 CCC	TTG TTG	294 ACG	348 GCT
CCC	GCT	ACG	CTC	CAA
ACC	GTC 	200	CAA	TTC
123 TCA 	AAT N	231 CCT	285 AAT 	339 TTG
GTC	GTC	GTT 	ACG	GGA
GCG 	AAC		GTC	
1114 GCT	168 AAG	222 GGT	276 AAT 	330 TGG
CTC	ATC	AAC	ATT	CAT
TTG	GAC	GTC	CGC	ATT
105 CCT	159 TTC	213 TCC	267 TTG	321 ACG
CTA 	AAG		ACC	ACA T
CTC	TAT	ATC	GAC	GCC
96 ACC 		204 TCT	258 GGT	312 CGT
ATT	CGC R	CGC	AAG K	CGT
AGC	GTC	CAG	AAC	ATG
87 TCT	141 GCC 	195 TTT 	249 GCC	303 AGT
CTT	GCT	200 200	ACG	CCT
ATG	TTT	GAT	ATC	GAC

F1G. 41

•								
402 CAA	α	456 TGG	3	510 GTC	>	564 AGC	S	618 AAG
ອວອ	4	ATG	Σ	TTG	J	909	4	GAA
ATT	H	ACC	E+	CCT	Д	GAT	Ω	CTA
393 CCT	<u> </u>	447 GGA	Ö	501 GGC	Ö	555 GAT	۵	609 GTT
TGC	၂ ပ	ACA	E					CCC
CAA	ļa	CAA	a	TTG	1	GAC	0	GCA
384 ACG	<u> </u>	438 GGC	ပ	492 GGA	ט	546 TAC	>	600 CCG
		၁၅၁	ĸ	GAT	Ω	၁၅၁	4	ACT
	<u> </u> [4	TTG	J	GTC	>	TCG	S	CAT
375 GCA	4	429 CCA	<u>d</u>	483 TAT	>	537 AAG		591 TAC
ည္သ	a.	ATC	Н	CAA	Ια	CAC	H	TGG III
ටු	ן ט	GAG	回	AGT	S	CCA	1 4	GAC
366 GAT	۵	420 TAC	; >	474 GCG	4	528 GAC	D	582 GAG
GAG	回	ACA	! ! [-	CTT	17	AAC	¦ z	CTT
GAC	۱۵	TAT	>	CAT	<u> </u>	CCA	1 4	ATG
357 GCC	4	411 TCC	S	465 GCC	4	519 GAT	۵	573 GTC
ACC	=		1	CAC	=	TAT	>	GTA V
ACT		AAT	Z	TAT	>	ATC	H	ACA

F I G. 4B

672 GGT 	726 GTA V	780 GCT:	834 GCC	888 608 1 1 0
TCG	TCA	TCT	GAG 	GCT
GAC	CGG	GCT	ATT	TAC
663 CCG 	717 CCC 	771 AAC 	825 GTC 	879 ATT
GTT	GTT	ATC 	ACT T	CAG
CCT	GCA	GTA >	CTG	TTC
654 TCT	708 CCC	762 CGC	816 AGT	870 AGC
CTC	GGT	TTG 	CAT	GAC
CTG	၁၁၁	000 R	GGA G	GTT
645 GCT	699 GTG	753 TAT 	807 GAA 	861 GCT
ACC	TAT	CGA	ATC	TTG
AAC	CGC	AAA	TCG	CC I
636 AAT	690 666 666	744 GGG	798 TTT 	852 CAG
ACT 	AAA	CGT	ACC	CAC
TCG	200	AAA	TYTT 	CTG
627 TTC 	681 AAT	735 GTA 	789 TCG 	843 ATC
ATG 	ATC 	AAC	5 5 5 5 5 5 5 5	5 5 5 5 5 5 5 5
CAA	CTT	ATC	ATC	GAT
			16/21	

F 1 G. 40

942 ATT	996 ACC	1050 ACG ACG	1104 GCG CTC 	1158 AAT CTT N L
TGG	CCC	ACG	GCG	AAT
TAC	GAC	DD A	CAT) T
933 AAC 	777G	1041 GAA E	.095 CTC 	149 TCC
GCC	AAC	1041 GCC GAA 	1095 AG AAC CTC (E N L	1149 GTT TCC
300 P - I	GCA	AAC	છ ા :	GAC
924 ACC 	978 AAT N	1032 GCG CCC 	1086 GTT GAA (140 GCA
CAA	ACC	GCG 	GTT	1140 CCC GCA P A
AAC	GGA	GGA	CTC	GCT
915 GCC 	969 GCC A	1023 TAC GAG 	1077 ACT GCA	1131 TCC
GAA E	GGA	TAC	ACT T	11 C GGC T
GTT 	GCA A	CAC	GGT))
906 ATC 	960 GTT 	1014 TTG	1068 F ATC	1122 CCG
GTC	ACC	GTA	GCT 	GCT
TCT 	ATG 	CCC	AGT	200
897 TAC 	951 CCA	1005 TTT 	.059 	1113 AAC CCT
290	GCA	GTC 	1059 A CAA GGC A	•
CAA	CGT	AAT	GAA 	ATC
			47/04	

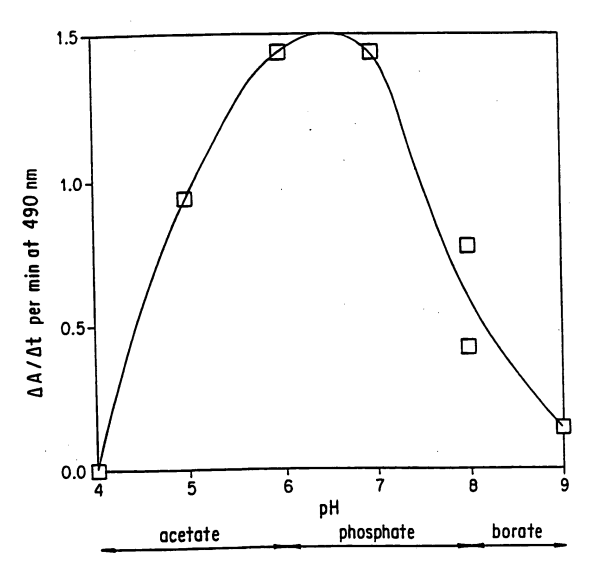
F 1 G. 40

0111	10 ft 1	061	ਵਾਹ:	& () i
1212 ATC	1266 GCG	1320 AAT	1374 CAC 	1428 AAC
1212 AAC ATC 	12665 AAT GCG 	CAC	1374 CTC CAC 	1428 GTC AAC
AAT N	AAC	CCA	CAT	TAT
1203 ACA TTT T	1257 TTG GCA 	1311 GTA TTG (1365 CCT ATC	1419 CCG AAC
ACA T	TTG	GTA >		္ ၁၂ ၂
TTC 	ATT 	ATC	CAC	ACC
1194 CTT AGG L. R	1248 TTG AAG	1302 CAC ACT 	1356 GCA GAC 	1410 GGT GGT
CTT	TTG	CAC	GCA	GGT
ATT 	CTC	GAG 	GGT	CTC
1185 GAT GGG D G	1239 CCC ACG	1293 CCA AAT 	1347 ACC GGA 	1401 AAA TCA
GAT D	CCC	CCA	ACC 1	AAA ×
GTT 	TTG	ACG	ATC	GTC V
1176 ACA T	1230 TCG	1284 TTC	1338 AAT	1392 ATC
AGC S	CCT	GAT	CTC	GAT D
CGC	GCT	GCC	GAG E	TTT
1167 . GGG	1221 GAG 	1275 GAC 	1329 ATC 	1383 GTG
ATT	1221 TAC GAG 	1275 AAT GAC 	GTT V	CAT
GCA	AAG K	AGC	AAA K	9 299
			18/21	

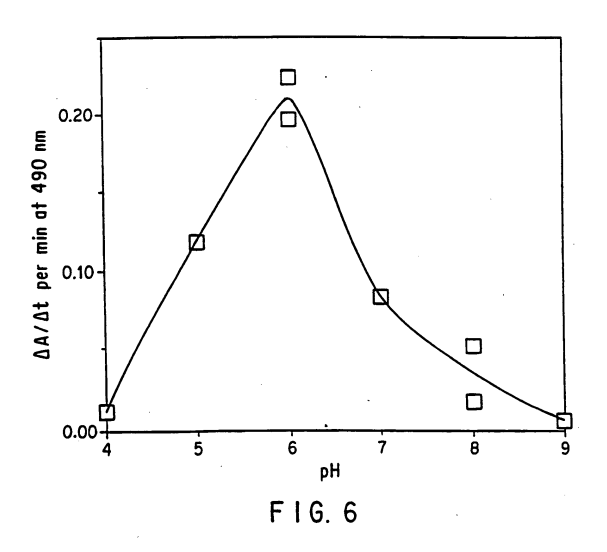
F16.4E

1482 TTC	1536 CAC TTG 	1590 CAG GGT 	1644 GCG	·
CGA F F F R	CAC	CAG	TAC	
CTC	TGG	CGC	AAG 	
1473 GTA	1527 GAC	1581 ATT	1635 CCC	
GTG 	1527 ATT GAC 	CAG	73C	·
GGT	CAC	AGC	CIC	
464 ACC	1518 TGC	1572 CCC P	1626 CAG	
36C	1518 CAC TGC 	1572 GCC CCC 	1626 AAC CAG 	
GGA 	GTT	GAG	TGG	
1455 GTC	TTT 	1563 GCC	1617 GCC	
CGT	TGG	TYT 	AAT 	T 3.
GTT 	CCA	GTC	AAC	CAG
1446 GTA V	1500 GGC G	CTT	1608 CCC	1662 TTG
GAC G	1500 CCA GGC P G	1554 GCA CTT 	16 CAG C	16(GAT TY D I
AGG	AAC	CTC	GTC)) 1
	1491 GAT D	1545 GGG	1599 TCG	1653 CCT
1437 CCA CGC P R	ACC	1545 GCT GGG 	1599 CAG TCG	CTT
CCG	AAG 	GAG	GTC	GCT
		19	9/21	

F16.4F



F I G. 5



21/21

A. CLASSIFICATION OF SUBJECT MATTER
IPC 6 C12N15/53 C12N9/02 C12N15/80 D21C5/00 A61K7/06
C12P7/22 C12N1/19 C09B69/10 //(C12N1/19, C12R1:66)

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols) IPC 6 C12N D21C A61K

Documentation searched other than minimum documentation to the extent that such documents are included in the fields executed

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

CHEMICAL ABSTRACTS, vol. 90, no. 19, 7 May 1979, Columbus, Ohio, US; abstract no. 147536w, BOLLAG J.M. ET AL. 'Characterization of an enzyme from Rhizoctonia praticola which polymerizes phenolic compounds.' page 213; see abstract & CAN. JOURNAL MICROBIOL., vol.25, no.2, 1979 pages 229 - 223	Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Y & CAN. JOURNAL MICROBIOL., 1,20-24, 29-41 39-41	X	7 May 1979, Columbus, Ohio, US; abstract no. 147536w, BOLLAG J.M. ET AL. 'Characterization of an enzyme from Rhizoctonia praticola which polymerizes phenolic compounds.' page 213;	14,43
-/	Y	& CAN. JOURNAL MICROBIOL., vol.25, no.2, 1979 pages 229 - 223	

Further documents are listed in the continuation of box C.	Y Patent family members are listed in annex.
*Special categories of cited documents: 'A' document defining the general state of the art which is not considered to be of particular relevance 'E' earlier document but published on or after the international filing date 'L' document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) 'O' document referring to an oral disclosure, use, exhibition or other means 'P' document published prior to the international filing date but later than the priority date claimed	"I" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art. "&" document member of the same patent family
Date of the actual completion of the international search	Date of mailing of the international search report
24 January 1995	23 . 02. 95
Name and mailing address of the ISA	Authorized officer
Buropean Patent Office, P.B. 5818 Patentiaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016	Delanghe, L

Vol.137, no.2, 1984 pages 89 - 96 Y WO,A,92 01046 (VALTION TEKNILLINEN TUTKIMUSKESKUS) 23 January 1992 see claims Y WO,A,92 16633 (NOVO NORDISK) 1 October 1992 see page 3; claims Y DE,A,30 37 992 (GESELLSCHAFT FUR BIOTECHNOLOGISCHE FORSCHUNG.) 19 August 1982 see claims Y EP,A,0 433 258 (ENSO-GUTZEIT OY) 19 June 1991 see claims Y EP,A,0 429 422 (ENSO GUTZEIT OY) 29 May 1991 see claims Y EP,A,0 408 803 (ENSO-GUTZEIT OY) 23 January 1991 see claims Y EP,A,0 060 467 (EISENSTEIN) 22 September 41	vol. 100, no. 19, us, Ohio, US; 20, 'The effect of pH on of syringic and the laccases of Dia and Trametes 1,20-24, 39-41
7 May 1984, Columbus, Ohio, US; abstract no. 152972q, LEONOWICZ A. ET AL. 'The effect of pH on the transformation of syringic and vanillic acids by the laccases of Rhizoctonia praticola and Trametes versicolor.' page 230; see abstract & ARCH.MICROBIOL., vol.137, no.2, 1984 pages 89 - 96 WO,A,92 01046 (VALTION TEKNILLINEN TUTKIMUSKESKUS) 23 January 1992 see claims WO,A,92 16633 (NOVO NORDISK) 1 October 1992 see page 3; claims WO,A,92 16633 (NOVO NORDISK) 1 October 1992 see page 3; claims PEP,A,0 433 258 (ENSO-GUTZEIT OY) 19 June 1981 see claims EP,A,0 429 422 (ENSO GUTZEIT OY) 29 May 1991 see claims EP,A,0 408 803 (ENSO-GUTZEIT OY) 23 January 1991 see claims EP,A,0 408 803 (ENSO-GUTZEIT OY) 23 January 1991 see claims EP,A,0 060 467 (EISENSTEIN) 22 September 41	2s, Ohio, US; 2q, The effect of pH on of syringic and the laccases of the lacc
ARCH.MICROBIOL., vol.137, no.2, 1984 pages 89 - 96 Y	39-41 TION TEKNILLINEN January 1992 O NORDISK) 1 October 21-24 SELLSCHAFT FUR FORSCHUNG.) 19 August SO-GUTZEIT OY) 19 June 40 SO GUTZEIT OY) 29 May 41 SENSTEIN) 22 September 41
TUTKIMUSKESKUS) 23 January 1992 see claims WO,A,92 16633 (NOVO NORDISK) 1 October 1992 see page 3; claims DE,A,30 37 992 (GESELLSCHAFT FUR BIOTECHNOLOGISCHE FORSCHUNG.) 19 August 1982 see claims PEP,A,0 433 258 (ENSO-GUTZEIT OY) 19 June 1991 see claims PEP,A,0 429 422 (ENSO GUTZEIT OY) 29 May 1991 see claims PEP,A,0 408 803 (ENSO-GUTZEIT OY) 23 January 1991 see claims PEP,A,0 060 467 (EISENSTEIN) 22 September 1982	January 1992 O NORDISK) 1 October SELLSCHAFT FUR FORSCHUNG.) 19 August SO-GUTZEIT OY) 19 June 40 SO GUTZEIT OY) 29 May 41 SO-GUTZEIT OY) 23 41 SENSTEIN) 22 September 41
1992 see page 3; claims Y DE,A,30 37 992 (GESELLSCHAFT FUR BIOTECHNOLOGISCHE FORSCHUNG.) 19 August 1982 see claims Y EP,A,0 433 258 (ENSO-GUTZEIT OY) 19 June 40 1991 see claims Y EP,A,0 429 422 (ENSO GUTZEIT OY) 29 May 41 1991 see claims Y EP,A,0 408 803 (ENSO-GUTZEIT OY) 23 41 January 1991 see claims Y EP,A,0 060 467 (EISENSTEIN) 22 September 41 1982	SELLSCHAFT FUR FORSCHUNG.) 19 August SO-GUTZEIT OY) 19 June 40 SO GUTZEIT OY) 29 May 41 SO-GUTZEIT OY) 23 41 SENSTEIN) 22 September 41
BIOTECHNOLOGISCHE FORSCHUNG.) 19 August 1982 see claims Y	FORSCHUNG.) 19 August SO-GUTZEIT OY) 19 June 40 SO GUTZEIT OY) 29 May 41 SO-GUTZEIT OY) 23 41 SENSTEIN) 22 September 41
1991 see claims P,A,O 429 422 (ENSO GUTZEIT OY) 29 May 1991 see claims P,A,O 408 803 (ENSO-GUTZEIT OY) 23 January 1991 see claims PP,A,O 060 467 (EISENSTEIN) 22 September 1982 41	SO GUTZEIT OY) 29 May 41 SO-GUTZEIT OY) 23 41 SENSTEIN) 22 September 41
1991 see claims Y EP,A,O 408 803 (ENSO-GUTZEIT OY) 23 January 1991 see claims Y EP,A,O 060 467 (EISENSTEIN) 22 September 1982 41	SO-GUTZEIT OY) 23 SENSTEIN) 22 September 41
January 1991 see claims Y EP,A,O 060 467 (EISENSTEIN) 22 September 41 1982	SENSTEIN) 22 September 41
1982	
see claims	RMA) 16 September 1992 42
X EP,A,O 504 005 (PERMA) 16 September 1992 42 see claims 42	

nformation on patent lamily inclines

PCT/US 94/10264

				·
Patent document sited in search report	Publication date	Patent memb		Publication date
WJ-A-9201046	23-01-92	NONE		
WO-A-9216633	01-10-92	AU-A- EP-A- JP-T-	1430992 0575462 6505873	21-10-92 29-12-93 07-07-94
DE-A-3037992	19-08-82	US-A-	4432921	21-02-84
EP-A-0433258	19-06-91	JP-A- NO-B-	3260188 174167	20-11-91 13-12-93
EP-A-0429422	29-05-91	CA-A- JP-A-	2030186 3174078	18-05-91 29-07-91
EP-A-0408803	23-01-91	DE-D- ES-T- JP-A- NO-B-	68912322 2061857 3130485 175105	24-02-94 16-12-94 04-06-91 24-05-94
EP-A-0060467	22-09-82	. DE-A- DE-A-	3110117 3128203	13-01-83 03-02-83
EP-A-0504005	16-09-92	FR-A- JP-A-	2673534 6172145	11-09-92 21-06-94